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THE UNIVERSITY OF ALBERTA

MEMBRANE ELECTROPHYSIOLOGICAL PROPERTIES OF DEVELOPING SKELETAL MUSCLE CELLS

by
Joy A. Steele

A THESIS

Department of Physiology

Fall, IOR4

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The undersigned certify that they have read, and recommend to the Faculty of Graduate Studies and Research, for acceptance, a thesis entitled MEMBRANE ELECTROPHYSIOLOGICAL PROPERTIES OF DEVELOPING SKELETAL MUSCLE CELLS submitted by Joy A. Steele in partial fulfilment of the requirements for the degree of Doctor of Philosophy.

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External Examiner

Date 4th September 1984

Dedication

To my husband, Gary and to my parents, Ann and Jim

Page 1

ABSTRACT

Isolated single fibers from the anterior (a. . and the posterior (p.l.d.) lattissimus/ dorsi muscles of bryonic and young chicks were used to study in vivo develop- ' ment of membrane electrical properties. Isolated fibers were obtained by an enzymatic dissociation procedure. Intracellular microelectrode recordings from isolated fibers and from fibers in intact muscles showed that the dissociation procedure did not significantly alter resting membrane potentials, input resistances or membrane time constants. The 14 day embryonic fibers of a.l.d. and p.l.d. did not have a measurable resting conductance to chloride. hatching, about 70% of the resting conductance in p.l.d. fibers was due to Cl. Membrane electrical properties were estimated from the analysis of voltage responses to intracellular injection of rectangular current pulses. At 14 days in ovo, membrane resistance (R_m) was approximately 20 kΩcm² and membrane capacitance (C_m) was 1-2 $\mu F/cm^2$ for both a.1.d. and p.l.d. The mean membrane length constants (λ) were for a.l.d. and 1.5 mm for p.l.d. The mean membrane time constants ($\tau_{\rm m}$) were 35.8 ms for a.l.d. and 25.3 ms p.l.d., the values of R_m , τ_m and λ decreased as development proceeded. For a.l.d., there was no change values by the time of hatching (21 days). decreases in the electrical constants for p.l.d. fibers were explained by the appearance of a resting conductance partly: during the last week of embryonic development.

Cultures of muscle cells derived from embryonic a.l.d. and p.l.d. muscles did not show fiber type differentiation in culture with respect to resting Cl conductance.

Membrane currents were recorded under voltage clamp from chick skeletal muscle with the whole-cell version of the patch clamp technique (Hamill et al., 1981). Muscle cells were grown in tissue culture in the presence of 10-* M colchicine which encouraged formation of sperical muscle cells (myoballs), Membrane Cl currents, which underlie a long duration, action potential, were studied in detail. Reversal potentials for the steady-state currents varied with Cl concentration in a manner predicted by the Nernst equation. Cl currents were blocked reversibly by SITS (1.0 mM) and SCN (10 mM) and irreversibly by DIDS (10 μ M). The currents activated slowly, following a delay, response to step depolarizations from a negative holding potential. Time constants for activation were obtained by fitting the rising phase of the currents with a single the opposite exponential and with another exponential of polarity to account for the delay. The major time constant was dependent on voltage and decreased from about 200 ms at -30 mV to about 50 ms at 30 mV. The Cl currents did not decline during maintained depolarization indicating lack of an inactivation mechanism. Following depolarizing pulses which activated the currents, currents which flowed during repolarization (tail currents) declined slowly completely. Tail currents were fitted with the sum

exponentials. One exponential component had an extremely long time constant on the order of seconds. The kinetics of the tail currents suggested that there must be a long-lived open state from which the channels returned very slowly to the closed state. Since three distinct time constants were found, there must be at least four kinetically distinct states: two closed states and two open states. Instantaneous current-voltage plots were linear indicating that the open channels behaved ohmically. Conductance values could then be calculated. Cl conductance showed a sigmoidal dependence on voltage. Conductance was low at potentials more negative than $-45\,$ mV, rose steeply between $-40\,$ and $0\,$ mV and leveled off above 0 mV. For potentials more negative than -25 mV, conductance changed e-fold for an 8 mV depolarization. Maximal conductance was 1.03 ± 0.7 mS/cm^2 (x Similar Cl currents were recorded from muscle cells that had developed in vivo, i.e. this Cl conductance mechanism is expressed during the course of normal developmont.

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Special thanks to Mark Poznansky for providing space, equipment, supplies, etc. and for allowing me to have my freedom. Many thanks to Ed Karpinski for expressing interest in the work and for assistance with the equipment. Also, thanks to Tessa Gordon for getting me interested in skeletal muscle development and to Doug Eaton for the loan of voltage clamp. Sue Faccio deserves thanks for typing the text of my thesis and Ken Burt's help with the figures is appreciated. Tella Findlay's help with all matter of things is also greatly appreciated. And to all of my companions in 'misery': Don Robertson, Damyanti Bhardwaj, Gary Simatos, Jean Gillespie, Janice Kuster, Glenn Wheeler, Davorka Krizaj Kapljic and Dave Reye, thanks for your friendships. Warm thanks to Sandra Czekarski for terrific company. And thanks to everyone elso in the Physiology Department who have been of aggighance.

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LIST OF NOTATION AND DEFINITIONS

a.l.d.	anterior latissimus dorsi
p.1.d.	posterior latissimus dorsi
TTX	tetrodotoxin
STX	saxitoxin
SITS	4-acetamino-4'-isothiocyano-2,2'-disulfonic acid stilbene
DIDS	4,4'-diisothiocyano-2,2'-disulfonic acid stilbene
m	Hodgkin-Huxley activation parameter
h	Hodgkin-Huxley inactivation parameter
n	Hodgkin-Huxley activation parameter
V _m	transmembrane potential expressed as potential of intracellular solution with respect that of extracellular solution (mV)
V _o	steady-state deviation of intracellular potential from resting potential recorded at the site of current injection (mV)
V _*	steady-state deviation of intracellular potential from resting potential recorded by an electrode other than the current passing électrode (mV)
x	interelectrode separation (µm)
đ	fiber diameter (µm)
λ	membrane length constant (mm)
R.,	input resistance (recorded potential/applied current) (M Ω)
۲ ,	intracellular resistance to axial flow of current along fiber (Ω/cm)
r _m	membrane resistance per unit length of fiber (Ωcm)
R _m	membranê resistance (Ωcm²)
R,	intracellular resistivity (Ωcm)
)	membrane time constant (ms)

 C_m membrane capacitance ($\mu F/cm^2$) G_m membrane conductance (mS/cm^2) R_* resistance in series with the cell membrane ($M\Omega$)

Other symbols are defined as they are introduced.

I. ION CHANNELS IN VERTEBRATE SKELETAL MUSCLE

The transmembrane ion movements underlying excitability in skeletal muscle fibers are controlled by voltage- and time-dependent membrane permeabilities. Protein molecules residing in the lipid bilayer appear to be responsible for these membrane properties. They form transmembrane pores which permit ion movements and are referred to as ion channels. Each type of ion channel has characteristic permeability, selectivity and kinetic properties (Armstrong, 1975; Hille, 1976; Neher & Stevens, 1977; Ulbricht, 1977; Armstrong, 1981; Hagiwara & Byerly, 1981).

Electrical excitation in adult skeletal muscle fibers involves voltage- and time-dependent changes of permeabilities to Na and K which allow a transient influx of Na ions followed by an efflux of K ions (Stefani & Chiarandini, 1982). As a result of these ion movements, an action potential is generated. In addition to Na and K' channels, a voltage-dependent Ca2+ channel has been described in several preparations (Hagiwara & Byerly, 1981; Stefani & Chiarandini, 1982). At the resting membrane potential, Cl and additional K channels are primarily responsible for resting membrane conductance (Adrian & Freygang, 1962; Palade & Barchi, 1977). Most of these K' channels pass inward current more easily than outward current and are known as inward rectifiers (Adrian, 1969).

Studies of ion channels in adult skeletal muscle have extensively utilized voltage clamp techniques. Since the

membrane permeabilities vary as a function of voltage and time, the ability to control voltage makes it possible to measure ionic currents directly. In addition, membrane voltage can be changed almost instantaneously, permitting separation of capacitative and ionic currents. A recent refinement of voltage clamp techniques has made it feasible to resolve current flowing through individual ion channels (Hamill, Marty, Neher, Sakmann & Sigworth, 1981).

The application of voltage clamp techniques to developing skeletal muscle fibers is technically difficult because of the small size and fragility of embryonic fibers. information concerning ion channels in of the developing fibers has been derived from studies of current injection. Several general points can be made however. The excitability properties change during the course of development (Spitzer, 1979). These developmental changes reflect differences in the types of channels that are expressed and differences in the densities of channels in the immature fibers. Other factors, such as differences in intracellular ion concentrations (McArdle, Michelson & D'Alonzo, 1980), may also be involved.

The following sections describe the properties of ion channels present in innervated and denervated adult muscle The same of the sa fibers and in developing muscle fibers. the same of the sa

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A. Ion Channels in Adult Skeletal Muscle

Sodium channels

ionic currents underlying the action potential in skeletal muscle were first described in frog muscle using a voltage clamp technique (Adrian, three-microelectrode 1970a). An early, transient inward Chandler & Hodgkin, current was identified as a Na current. It was blocked by tetrodotoxin (TTX), a specific blocker for Na channels (Narahashi, Moore & Scott, 1964), and reversed at a membrane potential close to the Na equilibrium potential. The time resolution for recording Na currents was greatly improved the double sucrose-gap by the introduction of (Ildefonse & Rougier, 1972) and the vaseline gap method * (Hille & Campbell, 1976), The development of current in response to depolarizing voltage steps was sigmoidal in shape. Currents were fit with a Hodgkin-Huxley kinetic model where $G_{Na} = G_{Na}^{max}m^{3}h$. The conductance to Na⁺ ions, G_{Na} , had a maximal value, G_{Na}^{max} , where nearly all of the channels were open. The third power of m was needed to describe the slow buildup of G following a step depolarization. The physical analogue of such a process is that three particles, voltage sensitive 'gates', must be 'near a certain area of the membrane at once in order for Na ions o cross. The probability of each gate being there is described by the parameter m. The activation variable, m, increased as potential was made more positive whereas the

inactivation variable, h, decreased.

Na conductance was found to be very voltage dependent. It rose steeply for small depolarizing pulses increasing e-fold for a 3.7 mV depolarization and leveled off above 0 mV. The membrane potential at which the conductance was half activated was around -50 mV and the time constant of activation (τ_m) at 0 mV ranged from 0.5 to 1.25 ms $(2-5^{\circ}$ C). The Na currents were found to inactivate, i.e. during a maintained depolarization the currents declined. Inactivation was voltage-dependent. The membrane potential for half inactivation was around -70 mV and the time constant of inactivation (τ_h) at 0 mV was about 1.5 ms. The information obtained from the voltage clamp data was used to numerically reconstruct the action potential (Adrian $et\ al.$, 1970a). Peak Na current density was about 4 mA/cm² and maximal Na conductance was 330 mS/cm².

Na* currents recorded from mammalian fast— and slow-twitch muscle fibers have been reported to be similar to those of frog muscle. Only minor modifications of m³h kinetics were necessary to fit Na* currents in rat muscle. (Pappone, 1980). The activation curve was shifted 10-15 mV in the hyperpolarizing direction from that found in frog muscle. This shift would increase the likelihood, of repetitive action potentials. Rates of activation at potentials more negative than -40 mV were significantly slower in rat than in frog. Values for maximal Na* conductance were similar in the two species. In rat, G_{Na}^{max} was 120 mS/cm² and

peak Na* current density was 2.5-4.5 mA/cm² (Adrian & Marshall, 1977; Duval & Léoty, 1978; Pappone, 1980). Slight differences between Na* currents in fast— and slow-twitch fibers have been noted (Duval & Léoty, 1980a). Peak Na* current occurred at less negative potentials and the reversal potential was about 7 mV more negative in slow-than in fast—twitch fibers. The difference in the reversal potential is consistant with the finding that the intracellular Na* concentration is higher in slow—than fast—twitch fibers (Sreter & Woo, 1963; Yonemura, 1967; Chutkow, 1973). Also, the density of Na* channels appears to be lower in slow—than in fast—twitch fibers based on differences in saxitoxin (STX) binding capacities (Hansen Bay & Strichartz, 1980).

Selectivity of Na* channels has been studied in frog (Campbell, 1976) and in rat (Pappone, 1980) muscle. The ratio of Na* to K' permeability (P_{Na}/P_K) was 0.048 for frog and 0.045 for rat. Permeability ratios for various cations were very similar to the results obtained by Hille (1975) for Na* channels in frog node to Ranvier. Hille characterized the ionic selectivity of the channels by systematically testing the permeability to an extensive list of small organic and metal cations. He proposed that a 3.1 x 5.1 Å oxygen-lined pore would account for the exclusion of all impermeant cations on simple geometric and chemical grounds.

Based on the density of TTX or saxitoxin (STX) binding sites in frog muscle, which ranged from 175 to 380 sites per μm^2 (Almers & Levison, 1975; Jaimovich, Venosa, Shrager & Horowicz, 1976; Ritchie & Rogart, 1977) and on maximal Natconductances of 50-330 mS/cm² (Ildefonse & Roy, 1972; Hille & Campbell, 1976), a single channel conductance of 1.5-15 pS was calculated. This range of values is close to the value of 18 pS determined from single channel records in cultured rat muscle (Sigworth & Neher, 1980).

Evidence has accumulated which indicates that channels also reside in the membranes of the transverse tubular system. This system of tubular membranes plays an important role in transmitting the electrical signals of the surface membrane into the central part of the muscle fiber to bring about a nearly synchronous initiation of contraction of myofibrils located at varying depths from the surface. The rad spread of mechanical activation has been reported to have a Q10 of 2 which is not easily explained by purely passive propagation of electrical signals along the tubular membranes (Gonzáles-Serratos, 1971). Spread mechanical activation was also reported to be less effective in Na deficient or TTX-containing solutions (Constantin, 1970). In addition, reduction of Na concentrations in the lumen of the transverse tubules or application of TTX to block regenerative activity of the tubular membranes has been shown to reduce twitch tension (Bezanilla, Caputo, Gonzáles Serratos & Venosa, 1972; Caputo & Dipolo, 1973;

Bastian & Nakajima, 1974). Detubulation of muscle fibers (glycerol-induced osmotic shock which interrupts the continuity between surface and tubular membranes) has been shown to reduce TTX binding capacity by about 50% (Jaimovich et al., 1976). Tubular Na currents have been recorded. They appeared as late, slow inward currents under voltage clamp (Hille & Campbell, 1976; Mandrino, 1977). Therefore, voltage-dependent Na currents in the membranes of the transverse tubular system contribute to the spread of depolarization along the tubules and are necessary for full mechanical activation.

Potassium Channels

Several types of K* channels have been described in frog skeletal muscle. 1) Delayed rectifier channels give rise to a fast outward current which is responsible for the repolarization phase of the action potential (Adrian et al., 1970b). The currents reach a maximum in about 100 ms at -30 mV (3°C). 2) Slow K channels generate slow currents that reach a maximum in about 3 s at 30 mV (3°C) (Adrian et al., 1970b). 3) The inward rectifier is responsible for a conductance increase when the membrane potential is shifted to more negative levels (Adrian & Freygang, 1962; Adrian et al., 1970b). 4) Ca² dependent K* channels, which open following increases in intracellular Ca² concentrations, have been postulated to exist in from muscle (Finh & Lüttgau, 1976; Nicola Ciri Sánchez & Stefoni, 1980).

Delayed rectifier: The presence of a clear repolarization phase of the action potential and of delayed currents in detubulated fibers demonstrate that delayed rectifier channels reside in the surface membrane (Gage & Eisenberg, 1969; Nicola Siri et al., 1980). There may also be delayed rectifier channels populating the transverse tubular membranes since the late afterdepolarization following a train of spikes was observed to be slower in larger diameter fibers (Kirsch, Nichols & Nakajima, 1977). The density of Kthannels has been estimated to be about 14 channels per μ m² which is 20 times less than that estimated for Nathannels (Almers & Levison, 1975).

Tetraethylammonium (TEA) has been shown to block delayed rectification in muscle as it does in nerve (Armstrong, 1975). However, the affinity of the binding sites in muscle was lower by an order of magnitude. TEA shifted the threshold and the activation curve to slightly more negative potentials and also slowed the rate of onset of the currents by about 80% (Stanfield, 1970b).

In frog muscle, the delayed outward K* currents could be fit with a Hodgkin-Huxley kinetic model (1952b) using n* kinetics, i.e. $G_K = G_K^{max}n^a$. The membrane potential at which activation was half maximal was around -45 mV and the time constant (τ_n) was 5-8 ms at 0 mV. The conductance increased e fold for a 3 mV depolarization. Maximal K* conductance ranged from 6-23 mS/cm² (Adrian et al., 1970a; Stanfield, 1970b. Stanfield, 1975; Almers, 1976). The instantaneous

current-voltage relationship was approximately linear with a mean equilibrium potential of -85 mV. The currents inactivated exponentially with a time constant of 0.5-1 s at 10 mV (20° C). The steady-state inactivation curve was less steep than that for Na current and the potential for half inactivation was approximately 20 mV more positive (Adrian et al 1970a).

Delayed rectifier currents is mammalian muscle have been reported to be similar to those in freq muscle (Pappone, 1980; Beam' & Donaldson, 1983a). Current density and time courses were similar in fact and slow twitch fibers of rat. However, slow-twitch fibers exhibited an additional larger slow K current which decayed ith a time constant about 10 times longer than that of the delayed current (Duval & Lécty, 1978; 1980a, b)

Slow potassium channels: In (ron muscle, 'ill current measurements suggested the precence of a slowly deviloring K' current that reached a maximum 'n about 3 s at 30 mV and then slowly inactivate to reach a final steady state. Teyel of about one third of the maximum amplitude (Advian 1 al., 1970b), slow K currents have been dire thy inscrided after blockade of the diayod rectifies with TEA in intertifiers (Stanfield, 1970a) and in the cut fiber preparation (Almers & Palade, 1981). The time course of the currents could be fitted with a Hodgkin moder (1957b) with a kinetics and the time course of the currents could be

the slow currents was about 10 mV more negative than that of the delayed current. This difference has been explained by postulating that the slow K channel is more selective for K than the delayed rectifier. Slow K current density was about one sixth of delayed rectifier current density.

slow twitch rat fibers, a prominent slow K* conductance has been reported to be present. In contrast to findings in frog muscle, the reversal potential for the delayed and slow outward currents was identical in detubulated fibers. In fast-twitch fibers, a much smaller slow component could be inferred from the tail currents. However, this component seemed to the result of K' accumulation in the lumen of the transverse tubular system as it was absent in detubulated fibers (Duval & Léoty, 1980a, b; Beam & Donaldson, 1983h) Therefore, it appears that fast-twitch muscle has only the fast delayed rectifier while slow twitch muscle has an additional slow K' current. function of the slow K' current is unknown, but its slow kinetics suggest that it may play a role in repetitive actimity. Slow twitch fibers do not show the repetitive firing in response to prolonged depolarization that fast-twitch fibers do (Duval & Légty, 1980a).

TEA and Zn² have been reported to block the delayed rectifier while having only a small effect on the slow current (Stanfield, 1975). In mammalian muscle, TEA blocked both currents while 4-aminopyridine blocked only the delayed the firm (Dival & Léoty, 1900b). Diethylpyrocarbonate, a

histidine reagent, has been reported to selectively block the slow current (cf Stefani & Chiarandini, 1982). The difference in pharmacology between the two currents supports the idea that the slow currents are carried by channels distinct from the delayed rectifier channels.

Inward rectifier The resting conductance of frog muscle shows inward-going or 'anomalous' rectification since it allows K' to move in across the membrane more easily than out (Adrian, 1969). Experiments on isolated single muscle fibers (Hodgkin & Horowicz, 1960; Adrian & Freygang, 1962) and on detubulated fibers (Eisenberg & Gage, 1969) strongly suggest that inward rectifier channels reside, for the most part, in the transverse tubular membranes. The channels show voltage- and time-dependent properties. In frog muscle, it has been shown that the conductance is activated by hyperpolarization (Hestrin, 1981; Leech & Stanfield, 1981). The conductance was found to increase e-fold for a 12 mV hyperpolarization. The time constant of activation versus voltage curve and the conductance versus voltage curve were found to shift to more positive voltages when the external K concentration was increased. No appreciable effect of intracellular K' concentration was found. Thus, the conductance mechanism depends on voltage and external R' concentration (Adrian & Freygang, 1962; Adrian et al., 1970b). Rectifia . dation of current flow through this K' channel seems to partly the result of voltage dependent gating (Hestrin, 1981) and partly the requit of 'instantaneous' rectification

of the channels since the instantaneous current-voltage relationship has been found to be nonlinear (Leech & Stanfield, 1981). The mechanism underlying rectification in individual channels has not been determined. The inward rectifier in fast— and slow-twitch mammalian fibers has similar properties to the inward rectifier in frog (Duval & Léoty, 1978; 1980a, b). The inward rectifier has been found to be blocked by TEA, Ba²⁺ and Sr²⁺, but not by Zn²⁺ (Stanfield, 1970a; Standen & Stanfield, 1978).

Inward rectifier currents recorded under normal ionic conditions (low external K*) have been observed to decline during maintained hyperpolarizing voltage steps. The decline has been demonstrated to be partly the result of K* depletion in the lumen of the transverse tubular system and partly the result of a fall in K* conductance for extremely large hyperpolarizing pulses (Adrian & Freygang, 1962; Adrian et al., 1970b; Almers, 1972). The fall in conductance for the extreme hyperpolarizing pulses (more negative than 150 mV) was shown to be the result of a potential dependent block by external Na* (Standen & Stanfield, 1979).

Moise analysis of inward rectifier currents in frog muscle has given a single channel conductance of about 9 ps (DeCoursey, Dempster & Hütter; 1984). Calculated channel density was 1 channel per μ m². Recordings of single channel currents, resulting from blocking and unblocking by Ba²⁺, in cultured rat muscle have also given a value of about 10 ps for single channel conductance (Ohmori, Voshida & Hagiwara,

1981).

In summary, two properties of the inward rectifier channels distinguish them from 'normal' delayed rectifier channels. First, conductance of the inward rectifier increases as membrane potential is made more negative. The opposite is true for delayed rectifier channels. Second, conductance of the inward rectifier channels depends on external K' concentration while the conductance of the delayed rectifier is independent of K' concentration. The molecular mechanism of this peculiar dependence on external K' concentration is not understood.

Calcium-dependent potassium channels: Ca2 - dependent K channels have been described in a variety of different cell types (for review, Kostyuk, 1984). The conductance depends on Ca2+ and on membrane potential. Studies of single channel events in cultured rat skeletal muscle have shown that elevation of intracellular Ca2+ levels increased the frequency and duration of the open state. A similar effect was produced by membrane potential with an e-fold increase in conductance for a 15 mV depolarization (Barrett, Magleby & Pallotta, 1982). Ca2+ dependent K channels from rabbit (Latorre, Vergara & Hidalgo, 1982) and from (Moczydlowski & Latorre, 1983) transverse tubular membranes have been studied in planar phospholipid bilayers. It was suggested that the channel requires two bound Ca2 ions for activation and that it is the binding steps which are voltage dependent while the opening and closing of

K'-selective pore is voltage-independent (Vergara, Moczydlowski & Latorre, 1984).

The single channel conductance has been reported to be 300 pS (Barrett et al., 1982; Latorre et al., 1982). The large conductance and the high Ca²⁺ and voltage sensitivity of the channel suggests that it is suited to resist depolarizations of the membrane potential which are accompanied by increases in intracellular Ca²⁺ concentrations. Thus, these channels may be a link between excitation-contraction coupling and membrane repolarization.

Calcium channels

Voltage clamp studies of frog muscle fibers have revealed slow inward currents carried by Ca2+ (Sánchez & 1978; Almers & Palade, 1981). Currents were Stefani, abolished by Co²⁺ and Cd²⁺, known blockers of Ca²⁺ channels (Hagiwara & Byerly, 1981), or by removal of external Ca^{2+} . Current-voltage relationships showed that currents were evident at -40 mV and reached a maximum at 0 mV. Maximum inward current was 80 $\mu A/cm^2$ and maximum conductance was 2-5 mS/cm². Ca² currents could be fit with a Hodgkin-Huxley kinetic model (1952b) using m³ kinetics. The membrane potential at which half-activation occurred was -39 mV and the time constant of activation (r_m) was 0.11 s at 0 mV. With maintained depolarization, currents declined completely. The time constant of inactivation was 1.1 s at 0 mV. However, it is entirely * clear what mechanisms underlie not

'inactivation' of the currents. It could be a voltage-dependent process (Cota, Nicola Siri & Stefani, 1983) or depletion of Ca²⁺ in the lumen of the transverse tubular system (Almers, Fink & Palade, 1981) or perhaps a Ca²⁺-dependent inactivation as seen in some nerve cells (Kostyuk, 1984).

Ca²⁺ channels have been reported to be located mainly in the tubular membranes (Potreau & Raymond, 1980; Almers et al., 1981). In detubulated fibers, a linear correlation was found between the degree of electrical continuity of the tubular with the surface membranes and the magnitude of the Ca²⁺ currents (Nicola Siri et al., 1980). The functional role of Ca²⁺ channels in muscle is not clear. Skeletal muscle fibers are well known to rely on internal stores of Ca²⁺ for regulating cytoplasmic Ca²⁺ concentrations. During a normal action potential, an insignificant amount of Ca²⁺ would enter the fiber through these channels to directly activate tension (Potreau & Raymond, 1980; Almers & Palade, 1981).

Chloride channels

cl channels make a significant contribution to the resting membrane conductance in avian, amphibian and mammalian muscle fibers. Resting Cl conductance has been reported to be (mS/cm²): 0.2 in frog (Hodgkin & Horowicz, 1959); 0.7 in Xenopus (Vaughan, McLarnon & Loo, 1980); 2 in rat (Palade & Barchi, 1977) and about 1 in chicken (Lebeda &

Albuquerque, 1975) muscle-fibers. In frog fibers, Cl channels appear to be located mainly in the surface membrane (Hodgkin & Horowicz, 1960; Eisenberg & Gage, 1969) while atleast 60% of the resting conductance to ${\rm Cl}^{-}$ (G $_{\rm Cl}$) appears to be located in the tubular membranes in mammalian fibers (Palade & Barchi, 1977; Dulhunty, 1979). An obvious advantage to having a large G_{Cl} in the tubular membranes would be that it would shunt the action potential and thus reduce ... outward K' current. This would then reduce the tendency for accumulate in the lumen of the transverse tubules. A casual relationship between K. accumulation and myotonic activity has been demonstrated in goats with a genetic (Adrian & Bryant, 1974). abnormally low $G_{\hbox{\footnotesize Cl}}$ has been postulated to produce some of symptoms of myotonia in humans (Bryant Morales-Aguilera, 1971).

G_{Cl} has been shown to depend markedly on external pH. The conductance has been observed to increase at alkaline pH and to decrease at acid pH. G_{Cl} and pH were related by a sigmoid curve with apparent pKs for the groups controlling G_{Cl} being about 7.0 in frog (Hutter & Warner, 1967; 1972) and 5.5 in mammalian fibers (Palade & Barchi, 1977). In frog fibers, voltage clamp records showed that G_{Cl} decreased as the fiber was hyperpolarized. For example, during a hyperpolarizing pulse of 85 mV from the holding potential, the current decayed with a half-time of 90 ms at pH.7.4. At alkaline pH (9.8), the decay was slower. In contrast, at

pH 5, current increased approximately exponentially throughout the pulse (Warner, 1972). The instantaneous current-voltage relationship was linear in the three pHs tested. However, the steady-state relationship suggested that the current reached a limiting value for large hyper-polarizations. According to Warner (1972), several features of G_{Cl} are consistant with the supposition that Cl ions combine with a carrier molecule to an extent determined by the external pH.

The Cl conductance in rat (Palade & Barchi, 1977) and in Xenopus (Vaughan et al., 1980) muscle has been shown to have similar properties to GCl in frog muscle. In Xenopus, it has been demonstrated that the voltage dependence of is similar in normally polarized and chronically depolarized fibers. This suggested that G_{Cl} is a function of the difference between resting membrane potential and membrane potential during a test pulse rather than a function of the absolute membrane potential. The findings in Xenopus and in rat fibers, have been explained by proposing the existance of aqueous pores for Cl (Palade & Barchi, 1977; Vaughan et al., 1980; Loo, McLarnon & Vaughan, 1981). To explain the observed deviation of the current-voltage relationship from that predicted from the constant field equation at alkaline pH; a blocking molety has been postulated. As the membrane potential becomes more negative, this particle would move to a site that must be occupied by Cl to permeate (Vaughan et al.; 1980). Gil in Xenopus fibers has been shown

blocked by the stilbene derivative, SITS (Vaughan & Fong, 1978).

B. Ion Channels in Slow Tonic Fibers

In addition to twitch fibers, there is another type of fiber in skeletal muscle: slow-tonic fibers. They occur either intermingled with twitch fibers in amphibians and mammals or as a pure slow muscle, the anterior latissimus dorsi, in birds (for review, Hess, 1970; Morgan & Proske, 1984). Slow-tonic fibers have multiple nerve endings, generate a slowly developing tension under repetitive stimulation and give rise to maintained or tonic tension when continuously depolarized with K or cholinergic drugs (Kuffler & Vaughan Williams, 1953b; Hess & Pilar, 1963; Page, 1969). This type of fiber is difficult to study electrophysic-logically because the fibers are scarce or have small diameters.

Sodium channels

In frog slow-tonic fibers, activatable Na channels appear to be absent (Burke & Ginsborg, 1956; Gilly & Hui, 1980). However, following denervation, these fibers were capable of generating propagating action potentials which were Na dependent and TTX sensitive (Miledi, Stefani & Steinbach, 1971). In contrast to frog fibers, avian fibers are capable of generating Na dependent action potentials which are TTX sensitive (Cullen, Harris, Marshall & Ward,

1975). The action potentials were reported to be smaller in amplitude and to have a slower rate of rise than the action potentials in twitch fibers. Mammalian slow-tonic fibers appear to be intermediate between amphibian and avian fibers, as they have been reported to show a graded response to depolarizing pulses. The response was blocked by TTX and disappeared in Na*-free solution (Bondi & Chiarandini, 1979; Chiarandini & Stefani, 1979).

Potassium channels

Delayed rectifier: The delayed rectifier has been reported to be present in frog, chicken and rat slow-tonic fibers (Burke & Ginsborg, 1956; Cullen et al., 1975; Bondi & Chiarandini, 1979; Chiarandini & Stefani, 1979). In froq fibers, maximum K' conductance was 0.5 mS/cm2 which is about ten times smaller than in twitch fibers. The steady-state conductance versus voltage curve was less steep than in twitch fibers with an e-fold change in conductance for a 15 mV depolarization. Also, the rates of activation were 2-4 times smaller than those of twitch fibers. Whether delayed rectifier inactivated during maintained depolarization was uncertain (Gilly & Hui, 1980). TEA blocked the delayed rectifier in mammalian and frog slow-tonic fibers (Bondi & Chiarandini, 1979; Gilly & Hui, 1980).

Slow potassium channel: So far, this channel has only been demonstrated in frog slow-tonic fibers (Gilly & Hui, 1980).

Calcium channel

Ca²⁺ channels appear to be present in slow-tonic fibers of the toad (Stefani & Uchitel, 1976). Studies of K⁺ and caffeine contractures in the anterior latissimus dorsi of chicks showed that the contractures were partially dependent on external Ca²⁺, suggesting that these fibers have Ca²⁺ channels (Page, 1969; Huerta & Stefani, 1981).

Resting conductances

fibers have been reported to have very low resting membrane conductances (mS/cm2): 0.008 in frog. (Stefani & Steinbach, 1969); 0.17 in chicken (Huerta, &. Stefani, 1981) and 0.25 in rat (Bondi & Chiarandini, 1979). The majority of the resting conductance has been reported to be the result of K' since resting conductance to Cl was not detectable. Current-voltage relationships have indicated that inward rectifier channels are present in chick and in rat (Chiarandini & Stefani, 1979; Huerta & Stefani, 1981), but not in frog (Stefani & Steinbach, 1969; Gilly & Hui, 1980) slow-tonic fibers. In rat, it has been reported that there is a small resting conductance to Na since Na free solution hyperpolarized the fibers by 10-15 mV (Bondi & Chiarandini, 1979).

C. Ion Channels in Denervated Skeletal Muscle

Sodium Channels

Deservation of rat skeletal muscle has been reported to cause a shift of activation and inactivation parameters of Na* currents by about 10 mV to more negative potentials (Pappone, 1980). The rates of activation and inactivation were not measurably affected. Since the resting membrane potential of denervated rat muscle has been reported to be more depolarized than normal (Redfern & Thesleff, 1971a), the shifts in activation could lead to significant Na* current at resting membrane potential. This is supported by the observation that the application of TTX increases the resting membrane potential of denervated, but not innervated fibers (Harris & Thesleff, 1971). The shift in activation may also explain the spontaneous, Na*-dependent, oscillations in membrane potential which cause fibrillation in denervated mammalian muscle (Purves & Sakmann, 1974).

The maximum Na conductance in denervated rat muscle has been shown to be similar to that in innervated muscle (Pappone, 1980). This is supported by the observation that saxitoxin (STX) binding capacity was only slightly reduced following denervation (Hansen Bay & Strichartz, 1980). In contrast to this, Barchi & Weigele (1979) have measured a 43% decrease in STX binding to purified sarcolemma of dener vated relative to innervated muscle.

It has been reported that the action potential becomes partially resistant to TTX following denervation (Redfern & Thesleff, 1971b; McArdle et al., 1980). In innervated muscle, TTX has been reported to block Na channels in a manner predicted by the binding of toxin to a single population of channels with a dissociation constant of 5 nm. Following denervation, a second population of Na channels with dissociation constant in the micromolar range appeared (Pappone, 1980). This second type of channel was responsible for 25-30% of the total Naticonductance. Thus, it appears that denervation causes, at most, a partial loss TTX-sensitive Na channels. It has been reported that denervated frog (Miledi et al., 1971) and chick (Lebeda; Warnick & Albuquerque, 1974) musgle remain normally sensitive to TTX. Therefore, denervation-induced TTX-insensitivity of Na channels is specific/to mammalian muscle.

Interestingly, following denervation of frog slow-tonic muscle, the fibers, which do not normally have Na channels, acquired to Na based action potential (Miledi et al., 1971). The amplitude and rate of rise of the action potential was smaller than in twitch fibers and reinnervation with a slow nerve repressed the action potentials. Thus, together with the above results, it appears that the expression of Na channels is at least partially influenced by innervation

Resting membrane conductances

Resting membrane resistance has been reported to increase following denervation of fast— and slow-twitch muscle fibers of rats (Albuquerque & McIsaac, 1970; Westgaard, 1975) and of chieffus (Lebeda & Albuquerque, 1975). This change has been determined to result from a decrease in resting Cl conductance (Lebeda & Albuquerque, 1975; Lorković & Tomanek, 1977). Interestingly, the increase in membrane resistance in rat muscle could be reversed by direct electrical stimulation of the muscle for 2 weeks beginning on the fifth day following denervation (Nestgaard, 1975). For denervated from muscle (over month), or hange in resting Cl conduct non-westfound But there is mostly decrease in resting Cl conduct non-westfound But there is mostly decrease in resting Cl conduct non-westfound But there is mostly

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A norther study of hapter in emitability during development of rational has pot been one. Action potentials has been recorded from diaghtage must be of 1. day rationages, administrational basis was retending the mond. & Miledia, 1969. Describing for the retending from diagrams and the bottom of a physical masses of the bottom of the retending for the respective of the physical masses of the bottom of the respective of the

TTX-insensitive Na channels.

A study comparing the action potentials in developing fast- and slow-twitch fibers showed that at 12 days of there were no differences between the two fiber types (McArdle et al., 1980). The characteristics of the action potential in slow-twitch fibers attained adult values by 18 days, while 50-60 days were required for the fast-twitch fibers. Differences between the two fiber types were apparent by 25 days. A quantitative study suggested that the basis for the differences between the two fiber types and between immature and adult fibers was the result of differences in intracellular Na* concentrations and maximal Na* conductance. Slow-twitch fibers were estimated to higher intracellular Na concentrations (29 mM) than fast-twitch (18 mM) and to have a lower Na conductance versus 34 mS/cm²). Young fibers were estimated to have very high Intracellular Na concentrations (45 mM for fast-twitch and 55 mM for slow twitch). The trend of high to low intracellular No concentrations during development was also found for another muscle of the rat (Atwood & Kwan, 1978) and for chick mustle (Barlow & Maneryy 1954):

The increase in maximum Na conductance during development probably reflected an increase in the density of Na hannels. Studies the number of STX binding sites per unit weight showed an increase from 2 to 21 fmol/mg wet weight from 2 to 25 days of age (Sherman & Catterall, 1982).

two phases. Suring the first phase, which was independent of continuing innervation, the STX receptor density increased to 47-57% of the adult level. After day 11, the second phase of development, which was dependent on continuing innervation, gave rise to the adult value of STX receptors.

Chick muscle

The development of excitability in chick muscle has been studied under current clamp (Kano, 1975). At 13 days in OVO, most of the fibers gave passive responses to depolarizing pulses, while a small percentage showed afterhyper-polarization, suggesting the presence of delayed rectification. At later stages, the first sign of regenerative activity was a long duration action potential subsequently shown to be produced by an inward Cl current (Fukuda, 1974; 1975; Fukuda et al., 1976a, b). By 19 days in ovo, a faster spike based on Na was present. Ca2 channels were also inferred to be present from the action of Ba2 on spike duration. Ba2 goes through Ca2 channels (Hagiwara & Byerly, 1981) and blocks K channels. After batching the Cl gotton potential disappears.

E. Ion Channels in Skeletal Muscle During in vitro
Development

Primary cultures of rat muscle

Developmental changes in action potentials have not been studied in primary cultures of rat muscle. However, the ratio of Na' to K' permeability (P_{Na}/P_{K}) was found to decrease from 0.4 at 3 days to 0.06 at 9 days of culture (Ritchie & Fambrough, 1975). This change in P_{Na}/P_{K} ratio was probably responsible for the increase in resting membrane potential (-24 to -51 mV) occurring over the same time period. Resting C1 conductance did not change with time in culture and remained well below adult values.

A novel C1 channel has been observed with single channel recording techniques (Blatz & Magleby, 1983). The single channel conductance was about 430 pS. The channel was often open at 0 mV. When open, voltage steps from 0 mV to either negative or positive potentials closed the channel after about 1 s. Returning the membrane potential to 0 mV caused the channels to reopen. In several trials, it was found that closed channels opened when the potential was stepped from ~60 to 10 mV for a few milliseconds and then back to ~60 mV. Once the channels opened, they closed within several seconds. If these channels have similar kinetics in intact cells, then they could possibly contribute to repolarization of the action potential.

. Clonal raticell line

Differentiation of electrical excitability studied in a clonal cell line of rat muscle (L6) (Kidokoro, 1973, 1975a, b; Land, Sastre & Podleski, 1973). Myoblasts were electrically excitable and produced small, brief Na* action potentials. After fusion, the multinucleate myotubes had overshooting action potentials. The initial spike was the result of Na influx and the plateau of the repolarizing phase was largely the result of Ca influx. These Na * channels may have differed from channels found in adult muscle since higher concentrations of TTX and STX were potentials required to block ion fluxes and action (Kidokoro, 1973; Sastre & Podleski, 1976; Lawrence & Catterall, 1981). Current-voltage relationships indicated that delayed rectification was present in myotubes but not in myoblasts. The time course of delayed rectification was thought to be extremely slow as the hyperpolarization following the action potential lasted 4-8 s. The slow kinetics of the delayed rectification may have contributed to the long duration of the action potentials in myotubes (about 100 ms at half height) (Kidokoro, 1973). Inward rectification was present in myotubes and the resting conductance to Cl was low (Kidokoro, 1975b). In summary, in this cell line, inward currents were expressed before out ward currents.

Primary cultures of chick muscle.

The excitability properties of chick skeletal muscle have been shown to change during development in tissue culture. After 2-3 days, inward and delayed rectifier conductances were present (Spector & Prives, 1977). Voltage-dependent Na*, Ca²* and Cl conductances appeared later, at days 4-5. (Kano, Shimada & Ishikawa, 1972; Kano & Shimada, 1973; Fukuda, 1974; Fukuda et al., 1976a). Claction potentials had extremely long durations (tens of seconds) and could only be elicited if the membrane potential was hyperpolarized below -60 mV or if Cl ions were injected intracellularly (Fūkuda, 1974; Spector & Prives, 1977).

The rate of rise of the Na* action potentials increased from 5 to 30 V/s during 4 to 14 days in culture (Kano & Yamamoto, 1977; Spector & Prives, 1977), but never reached adult values of around 112 V/s (Cullen et al., 1975). The increase in the rate of rise of the action potential appeared to be the result of an increase in the density of Na* channels since the binding capacity of TTX to muscle cell homogenates increased during 25 to 100 hours in culture (Frelin, Lombet, Vigne, Romey & Lazdunski, 1981). The authors did not find a change in the affinity for TTX. In another study, the appearance of STX and BTX (batrachotoxin) binding sites were found to follow different time courses (Strichartz, Bar-Sagi & Prives, 1983). These results suggested that the developmental expression of Na* channels

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was accompanied by a change in the pharmacological properties of the channel.

In cultured chick muscle, another novel channel has been found with single channel recording techniques (Kolb & Schwarze, 1984). This channel was cation selective, but lacked selectivity for one cation over another. The channel appeared to have voltage-dependent gating, but the kinetics were very complex.

F. Summary and Objectives

The surface membranes of mature skeletal muscle fibers are specialized to generate action potentials and to maintain an adequate transmembrane potential. It is now becoming apparent that these electrophysiological properties undergo changes during the course of muscle cell differentiation (Spitzer, 1979; Harvey, 1980). For example, the ionic dependence of the action potential changes during development. Observations such as this raise several questions. What is the exact nature and the time course of the changes in electrophysiological properties which occur during maturation of muscle? Are there changes common to other developing cell types, so that some general rules emerge? By what mechanisms do the changes occur? And do these changes play any role in the growth and maturation of muscle cells?

Muscle cells are attractive for the study of development of electrophysiological properties of membranes since a detailed knowledge about the mature fibers is available. Furthermore, a great deal of qualitative and quantitative information about membrane electrical properties can be obtained from intracellular microelectrode studies.

Unfortunately, there are technical problems which limit the study of electrical properties of developing muscles! At the embryonic stage of interest, the cells to be studied are generally small in size, making certain electrophysiological experiments. extremely difficult, if not impossible, to do. In addition, the fibers are often quite fragile and thus easily and rapidly damaged by microelectrode impalement. Consequently, very few studies and virtually no quantitative work has been done on developing skeletal muscle fibers.

One approach to these problems has been to prepare explant or primary cultures which can facilitate location, identification and experimental manipulation of the cells. Extremely valuable information can be obtained from study of development in vitro; however, care should be exercised in generalizing results to in vivo development since innervation affects the electrophysiological properties of muscle (for review see, Harris, 1974; Jolesz & Sreter, 1981). For example, it has been demonstrated that denervation of adult muscle leads to several changes: increased membrane resistance (Westgaard, 1975; Lebeda & Albuquerque, 1975; Lorkvic Tomanek, 1977) and the appearance of TTX-insensitive Na* channels (Pappone, 1980). Clonal cell lines have also been utilized (Kidokoro, 1975a, b), but they may demonstrate abnormal development.

In an effort to study in vivo development of vertebrate skeletal muscle, an enzymatic dissociation procedure previously applied to adult rat muscle (Bekoff & Betz, 1977a) was found to be applicable to embryonic chick muscle. The use of isolated single fibers greatly facilitated microstudies. Membrane electrical properties and electrode resting conductance to Cl were studied during embryonic development. Preliminary results on the voltage- and time-dependent ionic currents were also obtained using a single suction pipette voltage clamp. A detailed study of the voltage- and time-dependent Cl currents was done on muscle cells grown in tissue culture. The Cl currents underlie a long duration action potential and are expressed during the course of normal in vivo development. These action potentials appear to be involved in producing contractions (Spector & Prives, 1977).

II. MEMBRANE ELECTRICAL PROPERTIES OF DEVELOPING FAST-TWITCH AND SLOW-TONIC FIBERS OF THE CHICK

A. INTRODUCTION

The electrical properties (resistance, capacitance, time and length constants) of the membranes of skeletal muscle fibers determine their response to electrical stimulation. In birds and amphibians, slow-tonic and fast-twitch fiber markedly different membrane electrical types have properties (Adrian & Peachey, 1965; Fedde, 1969; Stefani & Steinbach, 1969; Gordon, Vrbová & Wilcock, 1981). Slow-tonic fibers receive multiple innervation and do not necessarily conduct action potentials (Ginsborg, 1960a; Ridge, 1971; Cullen et al., 1975; Gilly & Hui, 1980). The initiation of contraction in this fiber type involves the local spread of depolarization from the neuromuscular junctions. Slow-tonic fibers have higher membrane resistances and longer time and length constants than fast-twitch fibers which are innervated and rely on propagated action potentials to initiate contraction (Kuffler & Vaughan Williams, 1953a, b).

The membranes of slow-tonic and fast-twitch muscle fibers also differ in regard to resting membrane conductance to Cl. Fast fibers have a large resting conductance to Cl- (Hutter & Noble, 1960; Lebeda & Albuquerque, 1975) while slow fibers do not (Stefani & Steinbach, 1969; Huerta & Stefani, 1981). An abnormally low Cl conductance has been found to be partly responsible for the symptoms of myotonia

(Bryant & Morales-Aguilera, 1971; Adrian & Bryant, 1974). This finding supports the idea that the resting Cl conductance is important for the stability of the membrane potential especially during repetitive activity (Hutter & Noble, 1960; Rudel & Senges, 1972)

When do these distinctive properties of fast-twitch and slow-tonic muscle fibers arise during development? I have studied the anterior (a.l.d.) and the posterior (p.l.d.) latissimus dorsi muscles of the chick. A.l.d. is a multiply innervated, slow-tonic muscle and p.l.d. is a focally innervated, fast-twitch muscle (Ginsborg, 1960b). One week prior to, hatching, the embryonic fibers of both muscles had high membrane resistances (R_m), long time (r_m) and length (λ) constants and no detectable resting conductance to Cl (G_{Cl}). By the time of hatching, differences between a.l.d. and p.l.d. fibers became clear as a result of decreases in the magnitude of the electrical constants (R_m , r_m , λ) and the appearance of a substantial G_{Cl} in p.l.d. fibers.

B. METHODS

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Preparation

Latissimus dorsi muscles were dissected from white Leghorn chick embryos and young chicks (Figure 1) (Ginsborg, 1960a). The age of the embryo was determined from the number of days of incubation and the date of hatching (21 days) determined the post-hatched age. Single muscle fibers were

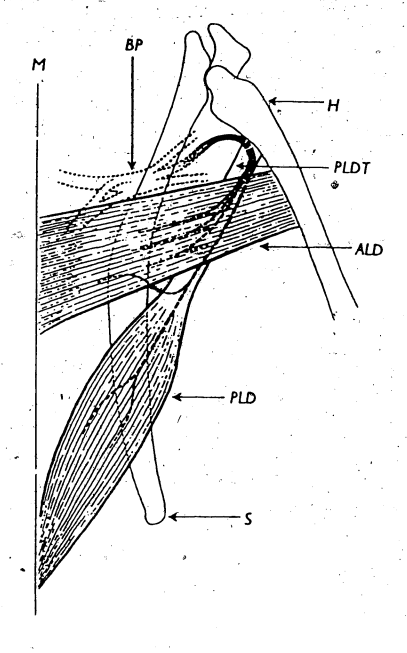


Figure 1. Diagram of a.l.d. and p.l.d. muscles. Dorsal view of the right latissimus dorsi muscle of the chick. ALD - anterior latissimus dorsi; PLD - posterior latissimus dorsi; PLDT - tendon of the p.l.d. muscle; M - midline; H - humerus; S - scapula; BP - brachial plexus. Adapted from Ginsborg (1960b).

isolated using an enzymatic and mechanical dissociation procedure (Bekoff & Betz, 1977a). Loose connective tissue was removed with fine forceps under a binocular dissecting microscope. Muscles were placed in Ca2+-free Eagle's minimum essential medium (MEM; Gibco, Calgary, AB) containing 0.3% collagenase (Sigma Chemical Co., Type I, St. Louis, MO). Muscles were incubated at 37° C in a water-saturated 5% CO₂ atmosphere until the fibers began to dissociate (approximately 1 hour for 14 day embryonic muscles and 2-3 hours for older muscles). After incubation, muscles were rinsed in Eagle's MEM containing 10% horse (Flow serum Inglewood, CA) and 5% chick embryo extract (CEE). CEE was made by pressing 11 day embryos through a 10 ml syringe. An equal volume of Earle's Balanced Salt Solution (BSS) wasadded. The composition of Earle's BSS is (gm/l): CaCl2, 0.2; KCl, 0.4; NaCl, 0.2; NaHCO₃, 2.2; MgSO₄, 0.2; NaH₂PO₄, 0.14; D-glucose, 1.0. The mixture was allowed to stand for 1 hour at room temperature. It was then centrifuged at 20,000 g for 10 minutes and the supernatant was removed and filtered (0.8 μ m). 5 ml aliquots were stored frozen until use. Following the rinsing step, muscles were mechanically dissociated release single muscle fibers. This was accomplished either by agitation with a jet of solution from Pasteur pipette or by trituration with a fire polished pipette. The mechanical dissociation step had to be done very gently to avoid damaging the fibers. Isolated fibers were placed in complete modic and hert in 35 mm tiesuo

culture dishes (Corning) in an incubator. Fibers were used the day they were isolated.

Experimental set-up

optics on an inverting microscope (Olympus). Fiber dimensions and interelectrode distance's were measured with a calibrated eyepiece micrometer. Solutions were made with glass distilled water and saturated with 95% O₂/5% CO₂. The external solution contained (mM): NaCl, 137; KCl, 5.4; CaCl₂, 1.8; MgCl₂, 0.8; dextrose, 10; HEPES (4-2-hydroxy-ethyl-1-piperazineethanesulfonic acid), 10. For Cl-free solutions, NaCl was replaced with NaCH₃SO₄ and KCl with KCH₃SO₄ (Pfaltz & Bauer, Stamford, CT) (Hutter & Noble, 1960) and MgCl₂ with MgSO₄, thereby replacing 98% of the Cl The pH was adjusted to 7.4. Experiments were done at room temperature.

Conventional techniques for intracellular recording and stimulation were used. Electrodes were conventional glass microelectrodes filled with 3 M KCl (20-60 M\Omega). Membrane potentials were recorded using WP Instrument electrometers (models 750 and M4-A). The current passing, electrode could be used to record voltage responses simultaneously. The resulting voltage drop across the electrode tip resistance was cancelled by balancing the active bridge circuit before impalement. Current was measured by a current-voltage converter placed in series with the bath ground. Signals were

displayed on a digital oscilloscope (Nicolet Instruments, Series 2090, Madison, WI) and stored on magnetic diskettes.

Leakage current from the electrometers was checked frequently and adjusted to as small a value as possible.

Digital voltmeters provided constant readout of membrane potential. Current pulse duration and magnitude were controlled with a digital pulse general of two truments was the day of the truments was checked frequently and adjusted to as small a value as possible.

Date collection and enalysis

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For estimation passive cable properties, polarizing rectangular current pulses were injected intracellularly. The magnitude of the current pulse was adjusted to give no more than a 15 mV response. Input resistances $(R_{i,n})$ and membrane time constants (r_m) were measured at site of current injection by the ourrent passing electrode. Membrane length constants (λ) were determined by inserting a Felimit electrode 100 cm away (distance limited by field of view) from the current pagaina Additional impalements for recording voltage responses at varying distancer from the current passing electrode were not feasible because of the lifficulty in obtaining multiple, stable impalements in the embryonic fibers. Since the current massing electrode was simultaneously recording voltage, care will taken to minimize any orrors from improper bridge balance or corrior, three fire primal path insight and the second of the second

Relatively low resistance electrodes were used and current pulses were as small as possible. Comparison of R_{1n} values in Table 2 (voltage measured with a separate electrode) with values in Table 3 (single microelectrode) show no significant differences (P < 0.1).

Passive cable properties were calculated from the equations for infinite cables with current injected in the middle of fiber length (Weidmann, 1952; Stefani & Steinbach, 1969). For all reported data, the distance from the current-passing electrode to the end of the fiber, divided by the length constant, was greater than 2, justifying use of the infinite cable equations (Stefani & Steinbach, 1969; Jack, Noble & Tsien, 1975). The length constant (λ) was calculated from $V_*/V_0 = \exp(-x/\lambda)$. V_0 is the steady-state deviation of intracellular potential from resting potential recorded at the site of current injection. V_x is the steady state deviation recorded at a distance, x, from the current passing electrode. Membrane resistance (R_m) was calculated from the following equations: $R_{ih} = V_0/I = r_i \lambda/2$; $r_m = r_1 \lambda^2$; $R_m = \pi dr_m$, r_1 and r_m are intracellular resistance and membrane resistance per unit length of fiber, respectively. d is fiber diameter. Intracellular resistivity (R.) was calculated by $R_i = r_i \pi d^2/4$. Membrane, capacutance, (C_m) equals t_m/P_m , t_m was taken as 84% of the time needed for the collage to reach with steady state value.

 $^{
m re}$ values reported in the text are from Student's t

C. RESULTS

Isolated fibers

Enzymatic dissociation procedures have been used to isolate single fibers from adult skeletal (Bekoff & Betz, 1977a) and smooth muscle (Singer & Walsh, 1980). No effects of enzyme treatment on resting membrane conductances were noted by Betz and Sakmann (1973) and only a slight decrease in resting membrane potential was observed by Bekoff and Betz (1977a).

Single muscle fibers were isolated from latissimus dotsi muscles of embryonic and young chicks. Visual inspection of the dissociated muscles at a magnification of 200x revealed dozens of single muscle fibers that appeared intact as well as fibers that had obviously been damaged. Damaged fibers had 'cytoplasmic blebs' and segments of them had undergone irreversible contracture. Many of the intact fibers settled to the bottom of the dish and adhered lightly to the substrate. They appeared cylindrical in shape and the banded pattern of sarcomeres was visible on fibers from 21 day embryos and older (Plate 1). Isolated fibers remained relaxed in standard external solution. Diameters of the isolated fibers were measured optically (see Ta' in 2): the

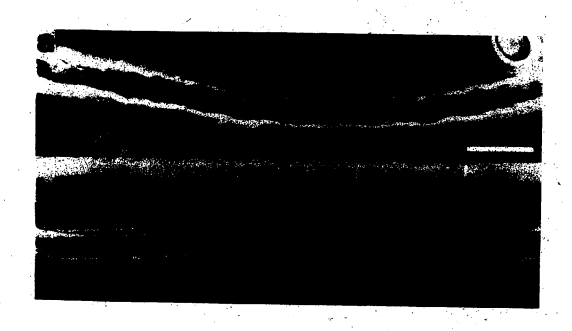


Plate 1. Light microscope photographs of single muscle fibers isolated from 14 A) and 21 B) embryonic posterior latissimus dorsi muscles of the chick. Calibration bar represents 20 μm

histological sections (Gordon, Purves & Vrbová, 1977).

Action potentials could be elicited by intracellular current injection and were often accompanied by visible contractions (Figure 2). At any one age, the amplitude, rate of rise and duration of spikes were quite variable from fiber to fiber. This variability probably arose because developmental changes in the action potential were occurring and the fibers were not developing synchronously (Kano, 1975).

Fibers that attached to the substrate along most of their length could be maintained in tissue culture for at least one week. During that time, the morphology of the fibers changed. The fibers sent out branches and began to resemble muscle fibers that have developed from myocytes in vitro. The cultured fibers exhibited spontaneous contractions and some began to fibrillate after several days in culture. Bekoff and Betz (1977b) reported similar changes for isolated adult rat fibers which had been maintained under tissue culture conditions.

This particular dissociation procedure was suitable for both a.l.d. and p.l.d. muscles dissected from embryos. It was not possible to isolated viable a.l.d. fibers from chicks older than several days primarily because of extensive amounts of connective tissue. The procedure worked well for p.l.d. muscles up to two weeks after hatching. After that age, enzymatic digestion of the extracellular material was fairly complete but the fibers were too long to

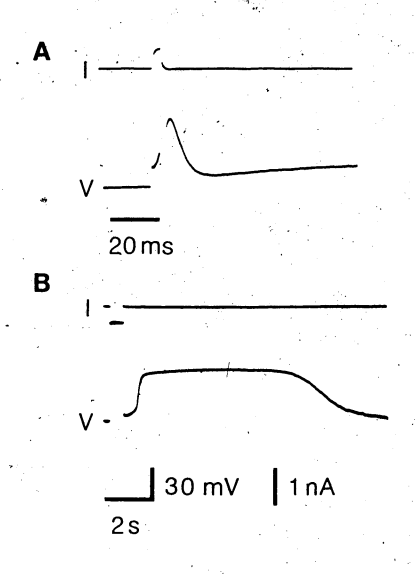


Figure 2. Action potentials elicited by current injection from a 21 day p.l.d. fiber. Resting potentials were -58 (A) and -65 mV (B). Potential at the top of the plateau (B) was -18 mV. Fibers were bathed in normal external solution.

successfully extricate from the network of blood vessels.

To assess the condition of the isolated fibers, values for resting membrane potential (V_m) , input resistance (R_{in}) and membrane time constant (τ_m) were obtained and compared to values obtained from fibers in intact muscles. Action potentials were not used as a basis of comparison because of the variability amongst fibers (cf Kano, 1975). Comparison of the values from isolated and intact fibers (Table 1) showed that the dissociation procedure did not have a significant effect on these properties.

Chloride conductance during development

Resting membrane conductance to $\operatorname{Cl}^-(G_{C_1})$ was estimated by measuring R_{in} and r_{m} values of single fibers in normal external solution and in a solution in which Cl was replaced by an impermeant ion (methylsulphate) (Table 2). No significant differences between the two solutions were found for either p.l.d. or a.l.d. muscles at 14 days or for a.l.d. at 21 days (P > 0.2), indicating little or no participation of Cl in the resting membrane conductance. For 21 day p.l.d., R_{in} and τ_m were much higher in the Cl -free solution, indicating a substantial resting conductance to Cl. The contribution of G_{Cl} to the total resting membrane conductance (G_m) was estimated assuming that membrane capacitance (C_m) remained unchanged. Since au_m is proportional to the inverse of G_m and since G_m is the sum of the conductances to Cl and the remaining ions

Table 1. Electrical properties of intact and isolated a.l.d. and p.l.d. fibers. Recordings were made from fibers in intact muscles and from isolated fibers. Muscles were from 21 day embryos. A single microelectrode was used to simultaneously inject current pulses and record voltage responses. All recordings were made in normal external solution. The number in parenthesis gives the number of fibers.

* 0.1 > P > 0.05; all other P > 0.2

Muscle	Condition	V _m (mV)	Rin (MΩ)	τ _m (mse _c)	
p.1.d.	Intact (23)	-64.2 ± 6.8	19.7 ± 6.3	18.3 ± 6.7	
	Isolated (36)	-59.5 ± 7.2*	20.2 ± 9.2	15.7 ± 5.5*	
a.l.d.	Intact (37)	-62.7 ± 6.5	14.7 ± 4.2	30.9 ± 11.0	
	Isolated (24)	-61.8 ± 5.9	15.5 ± 5.5	33.7 ± 20.6	

Table 2. Resting chloride conductance of 14 and 21 day a.l.d. and p.l.d. fibers. Recordings were made from isolated fibers with one electrode to inject current and another electrode inserted nearby to record voltage responses. Fibers were equilibrated for 30 min to 1 hr in a solution with 98% of C1 replaced with CH₃SO₄. Fibers were then returned to normal solution and examined. The number in parenthesis is the number of fibers examined.

Muscle	Age (Days)	Solution	Rin (MΩ)	τm (msec)	
p.l.d.	14	Normal (20)	44.2 ± 22.0	30.2 ± 13.2	
. (14	Cl free (22)	40.2 ± 18.4	28.9 ± 12.1	
	21	Normal (31)	18.3 ± 10.4	15.4 ± 5.9	
4	21	Cl free (25)	41.5 ± 11.6	53.9 ± 7.8	
a.l.d.	14	Normal (20)	25.9 ± 9.2	33.7 ± 15.2	
	1.4	Cl free (21)	22.2 ± 7.4	28.9 ± 11.9 -	
	21	Normal (25)	15.0 ± 4.7	36.0 ± 13.0	
	21	C1 free (24)	15.5 ± 4.8	38.4 ± 8.4	

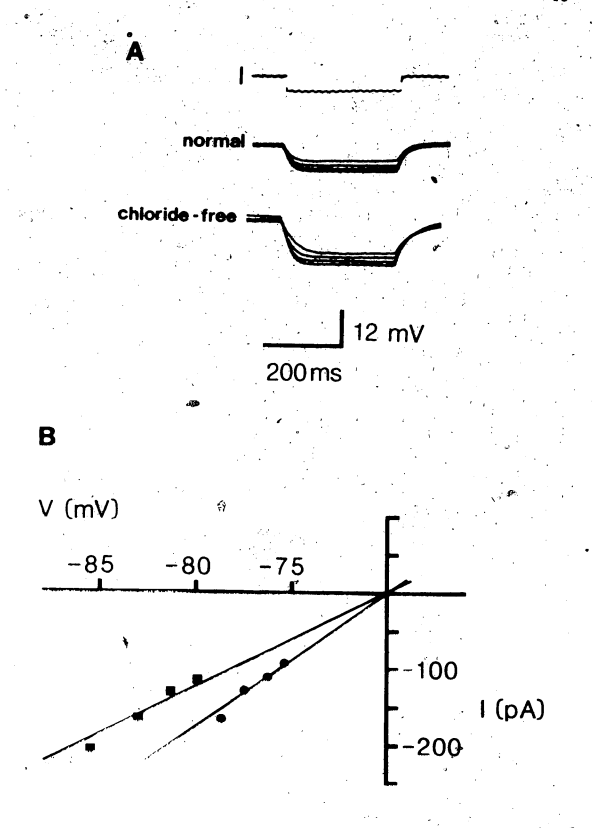
 $(G_m = G_{Cl} + G_{lon})$ (Eisenberg & Gage, 1969), it was calculated that G_{lon} constituted about 70% of the total resting membrane conductance in 21 day p.l.d. fibers. Thus, it was not until the last week of embryonic life that this type-specific property was expressed p.l.d. fibers.

Myotubes were grown in tissue culture (see Methods, Chapter III) from myocytes obtained from 15 day a.l.d. or p.l.d. muscles. Resting G_{Cl} was determined in cultures 5-10 days old. Figure 3 shows the results from a myotube where $R_{i,h}$ increased from 56 to 80 $M\Omega$ when the solution was changed to Cl -free. This was the largest increase observed. In general, R., values only increased 5-20% when solution was changed to Charge. For p.l.d. cultures, Rin values were 75 \pm 45 M Ω in normal and 90 \pm 49 M Ω (% n = '11) in Cl -free solution./ For a.l.d. cultures, Ri values were 60 \pm 25 M Ω in normal and 66 \pm 22 M Ω (% \pm n = 11) in Cl -free solutions. Rin values showed considerable variability and were not significantly different between normal and Cl -free solutions for either culture (P > 0.2). Thus, fiber type differentiation with respect to this property did not occur in culture.

Passive cable properties during development

Values for cable properties were obtained by analysing voltage responses to injected current pulses. Voltage was recorded at two separate sites (Figure 4) and only occasionally could two stable impalements be made simultaneously.

Figure 3. Resting chloride conductance of cultured p.l.d. muscle. Myocytes were obtained from 15 day p.l.d. muscles and grown in tissue culture for 5 days. A single microelectrode was used to inject rectangular current pulses and to record voltage responses (A). Responses were first recorded in normal (149 mM Cl⁻) and then in Cl⁻-free (3 mM Cl⁻) solution while continuously recording from the same myotube. Cl⁻ was replaced with an impermeant ion, methylsulphate. Current-voltage plots of the data (B) gave an input resistance of 56 MΩ in normal solution (circles) and of 80 MΩ in Cl free solution (squares).



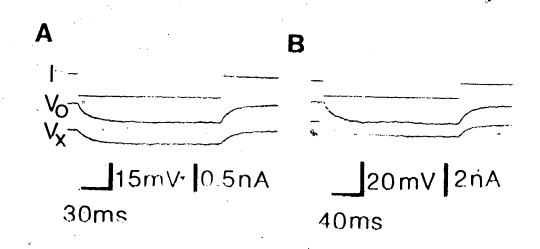


Figure 4. Passive electrical responses of isolated a.l.d. and p.l.d. fibers. Voltage responses were recorded from 21 day p.l.d. (A) and a.l.d. (B) in normal external solution. I is applied current pulse injected in the middle of the fiber length. Voltage response recorded by the current injection electrode. ". is the voltage response recorded by the current injection electrode.". is the voltage response of isolated a.l.d. by a second electrode 460 (A) and 620 μm (B)

The measured and calculated values are presented in Table 3. At the earliest stage studied, 14 days' incubation, the time and length constants for a.l.d. and p.led. fibers were and the means for the two muscles differed only slightly (P < 0.2). R_m values were fairly high (20 $k\Omega cm^2$) and the means for the two muscles were not different (P > 0.1). C_m values (1-2 $\mu F/cm^2$) were in line with what would be expected for small diameter fibers (Hodgkin & Nakajima, 1972). As development of p.l.d. proceeded, the values of R_m , τ_m and λ that by the time of hatching, differences decreased, S₽ between a.l.d. and p.l.d. were apparent (P < 0.001). For fibers, R_{m} did not differ at the two ages studied (P > 0.1). For p.l.d., Rm decreased further after hatching. fibers were not followed after hatching because the fibers could not be isolated. The values of V_{m} reported here. closely agree with '' alar of Bennett and Pettigrew (1974) and (1975).

P PISCUSSION

Before the mechanisms governing differentiation of muscle fibers into various fiber types can be understood, it is important to know when type specific properties are expressed during development. Knowledge of the timetable of developmental changes may help to elucidate causal relationships. 'n the present study of developing fast twitch (p.1.d.) and slow toni (a.1.d.) muscles of the chick, a

Actrical constants calculated see Methods). The number of parenthesis is the number of fibers examined. angle libers the day they were solated. Rectangular current pulses were anjected into the middle of the ber and the voltage response recorded for the determination of \Re_{10} and τ_{m_0} . A voltage-sensing electrode able ?, Electrical constants of a.i.d. and p.i.d. libers during development. Recordings were made from as aserted 400-600 um away for the setermination of A. Fiber diameters were measured optically, and

Ri (Acm)	169 ±49	103 155	.26 £45	144	30	:55 ±34
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π xscm²)	.2.9 ±4.3	2. 51 5. 53	4 60	2 1:15	20.2	7.8 ±5.i
(MOcm)	410	7	1.0 4.02	7.8	3.6	i. i. i.
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presumptive fast-twitch muscle at an early stage of development (Table 2). By the time of hatching, the percentage of total membrane conductance due to Cl had reached adult levels. Expression of this property, which is characteristic of adult p.l.d. fibers (Lebeda & Albuquerque, 1975; Huerta & Stefani, 1981), followed the formation of functional end-plates (Gordon, Perry, Tuffery & Vrbová, 1974; Atsumi, 1977). In contrast, G_{Cl} in developing a.l.d. muscles was negl ble at both ages studied (Table 2). These results fit well with those of Huerta and Stefahi (1981). They found that at three weeks of age, about 70% of the resting conductance of p.1.4: fibers was due to Cl-, while a.1.d. and the have a measurable Gol. An increase in the magnitude of Goodwine post-natal development of mammalian fast muscle, but no change for a slow muscle; has been reported ti Parnhach, Brown and Barchi (1978).

These results raise the possibility that innervation is required for full expression and/or maintenance of G_{Cl} . The court but on of G_{Cl} to the total resting membrane conductions ass found to be low (5-20%) for chick muscle cells grown in tiscus culture. This appears to be a general finding as G_{Cl} has been found to be low in aneural cultures of rat (hitchie & Fambrough, 1975), of rat cell line (hidek or loss) and of human (Merickel, Gray, Chauvin & Appel 1981) skelmal muscle Also, the slight increase in membrane sistance following denorvation (Albuquerque & Ibestol 1969, Usetoned, 1975) has been attributed to a

decrease in $G_{\rm Cl}$ (Camerino & Bryant, 1976). It would be useful to understand the factors controlling expression of $G_{\rm Cl}$ as its magnitude is abnormally low in myotonic muscle (Bryant & Morales-Aguilera, 1971) and in dystrophic mouse muscle (Farnbach et al., 1978).

The lack of clear differences in resting $G_{\rm Cl}$ between muscle cell cultures derived from a.l.d. or p.l.d. muscles suggests that fiber type differentiation does not occur in culture. This conclusion has also been reached from studies of metabolic enzymes, which show type-specific levels of activity, in chick and rat cultures (Askanas, Hee & Milhorat, 1972; Askanas, Shafiq & Milhorat, 1972; Askanas & Hee, 1973). However, slight differences in favor of expression of fiber type in culture have been obtained (Nougues & Bacou, 1977; Bacou & Nougues 1980). In general though, it appears that the instructional mignal(s) directing fiber type differentiation is(are) lacking in the timesue culture environment.

The absolute values obtained for the cable properties of developing slow-tonic (a.l.d.) and fast twitch (p.l.d.) fibers (Table 3) were higher than previously reported for fibers in intact muscle (Gordon et al., 1977). Since the input resistances and the resting membrane potentials were also much higher, the higher values probably reflect more stable impalements. Also, the method used to estimate the cable properties was more direct than that used by Gordon et al. (1977). They calculated the action of their input

resistances and a mean value of fiber diameter obtained from histological sections.

One week prior to hatching, future slow-tonic and fast-twitch fibers had relatively high membrane resistances, long time and length constants (Table 3) and no detectable resting Cl conductance (Table 2). Thus, the properties of embryonic fibers of both muscles resembled more closely properties of adult slow-tonic rather than fast-twitch (Fedde, 1969; Gordon et al., 1981). By the time of hatching (21 days), R_{max} , r_m and λ of p.1.d. fibers, had decreased so that the cable properties of the a.l.d. and p.l.d. fibers were clearly distinguishable. In both fiber R_{m} would be expected to decrease and C_{m} to increase types, as the fibers grow in diameter simply because the ratio of transverse tubular to surface' membrane is increasing (cf Hodgkin & Nakajima, 1972). For the developing p.l.d. fibers, however, the decreases in the electrical constants were mostly attributable appearance of a substantial to the resting conductance to Cl during the last week of embryonic down opment

III. VOLTAGE- AND TIME-DEPENDENT CHLORIDE CURRENTS IN EMBRYONIC CHICK SKELETAL MUSCLE

A. INTRODUCTION

Membrane chloride conductances have been found in a wide variety of cell types and in a variety of species, ranging from E. coli and algae to higher organisms. Utilizing current clamp techniques, Cl -dependent action potentials have been observed in certain plant cells (Gaffey Mullins, 1958; Mullins, 1962), in electroplax membranes of some fishes (Bennett, 1961; Hille, Bennett & Grundfest, 1965) and in frog eggs (Ito, 1972). With voltage clamp techniques, Cl currents responsible for repolarizing a Na -based action potential have been described in frog eggs (Schlichter, 1983). In Aplysia neurones, Cl currents actihyperpolarization by have (Chenoy Marchais, 1982). Voltage and time-dependent Cl currents have also been described in frog (Hutter & Warner, 1972: Warner, 1972) and in Xenopus (Vaughan et al., 1980; Loo et al., 1981) skeletal muscle. With single channel recording techniques, Blatz and Magleby (1983) described a voltage-dependent Cl channel with a single channel conducfance of 430 pS in cultured rat muscle. The properties of some Cl selective channels have been studied by incorporating the channels into planar phospholipid bilayers. Cl. channels with single channel conductances of 16 and 55 ps have been found in electroplax membranes of Torpedo (marine

ray) (White & Miller, 1979, 1981; Hanke & Miller, 1983) and in heart sarcolemma of calves (Coronado & Latorre, 1982), respectively. A voltage-dependent anion-selective channel (VDAC) from mitochondrial outer membrane with a single channel conductance of about 500 pS has been described (Schein, Colombini & Finkelstein, 1976; Colombini, 1979; Doring & Colombini, 1984). Further, ion tracer techniques have been utilized to study a Cl- transport system in peripheral blood lymphocytes activated by increases in cell volume (Sarkadi, Mach & Rothstein, 1984) and to study a Cl- permeability system in red blood cells which is mediated by band 3 protein (Cabantchik, Knauf & Rothstein, 1978).

Voltage-dependent anion conductance systems have not been as extensively studied as cation conductance systems. In this chapter, I describe the properties of a time-variant, voltage-dependent Cl conductance found in embryonic chick skeletal muscle. The conductance underlies a long duration action potential (Fukuda, 1974; Fukuda et al., 1976a). The action potential can be elicited from the muscle fibers during the last week of embryonic development (Kano, 1975).

Preliminary voltage clamp studies have been done using tissue cultured muscle (Fukuda, 1975; Fukuda et al., 1976b). In the present study, the surface membrane of small diameter myoballs was voltage-clamped with a single suction pipette (Hamill et al., 1981). The data showed that the Cl-currents activated slowly in response to depolarizing voltage steps

and did not inactivate during maintained depolarization. Upon repolarization to negative holding potentials, the currents recovered extremely slowly from activation. Cl-currents were blocked by stilbene derivatives which are known to block other Cl-conductances (Knauf & Rothstein, 1971; Vaughan & Fong, 1978). The Cl-currents were also found in muscle cells that had developed *in vivo*, showing that this Cl-conductance is indeed expressed during the course of normal development.

B. METHODS

Preparation of myoballs

Small diameter (10-20 μ m) myoballs (spherical cells) were grown in tissue culture. Under sterile conditions, pectoral muscles were removed from 11 day white Leghorn chick embryos and placed in Earle's balanced salt solution (BSS). The muscle was minced into small fragments (1 and then triturated with a fire-polished Pasteur pipette to release single cells. The solution was left for 5-10 minutes to allow debris to settle. The top layer was removed and filtered through lens paper. The cells were gently centrifuged in a clinical centrifuge and resuspended in BSS. This was repeated three times, and after the last centrifugation the cells were resuspended in complete media. Complete media consisted of: Eagle's minimum essential media (MEM), 10% horse serum (Flow Labs,

Inglewood, CA), 5% chick embryo extract (see Methods, Chapter II), penicillin (100 U/ml) and gentamycin (200 μ g/ml). Cell, density was determined with a hemocytometer. Cells were plated at 0.8 x 10^6 cells/ml in 35 mm Corning tissue culture dishes. Plates were placed in a water-saturated 95% $O_2/5\%$ CO_2 atmosphere and maintained at 37°C. On day two of culture, the media was changed to one containing 10-8 M colchicine (Sigma, Type II, St. Louis, Colchicine encouraged the muscle cells to form MO). spherical myoballs (Plate 2), but did not appear to have any detrimental effects on membrane or cellular properties (Fukuda et al., 1976a). Media was changed every three days.

Myoballs were also obtained from muscle fibers that had developed in vivo. Isolated fibers were dissociated from intercostal muscles of 14-21 day embryos according to the procedure outlined in Chapter II (Methods). These muscles were used because the fibers are only several millimeters in length. Isolated fibers often broke into small lengths with more rigorous trituration. Fragmented fibers often sealed and after several hours of incubation spontaneously formed myoballs of various diameters (20-70 μ m).

Experimental set-up

Membrane currents were recorded under voltage clamp with the whole-cell recording technique of Hamill et al. (1981). Briefly, suction was applied to a wide-tipped micropipette to establish a tight seal, in the giga-ohm range,

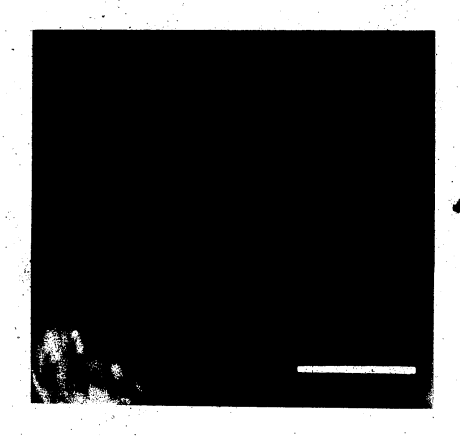
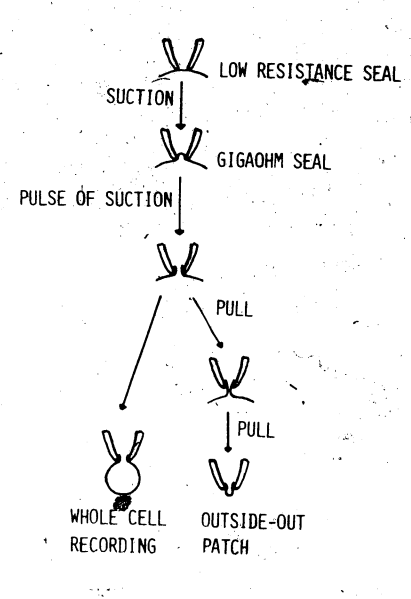


Plate 2. Light microscope photograph of a myoball grown in the presence of 10^{-8} M colchicine. Myocytes were obtained from pectoral muscles of 11 day chick embryos. Calibration bar represents 20 μm .

between pipette tip and cell membrane. The membrane patch spanning the pipette tip was disrupted by additional suction so that the suction pipette became an intracellular electrode (Figure 5). This procedure provided a low resistance pathway to the intracellular space.

Fabrication of suction pipettes: Suction pipettes were pulled according to the methods of Hamill et al. (1981) from microstar capillary tubing with an O.D. (Radnoti Glass Technology, Inc., Monrovia, CA). Pipettes were pulled in two stages on a vertical Narishige microelectrode puller fitted with a four-turn coil of nichrome wire (1 mm diameter). In the first pre-pull, the capillary tubing was thinned over a 8-10 mm length to a minimum diameter of about 200 μm . The tubing was then recentered with respect to the heating coil and in the second pull, the thinned part broke to produce two pipettes. This two stage pull produced pipettes with short shanks and tips with steep tapers. No magnetic pull was employed. The heat setting was critical (11-11.5 A) as slight variation produced large variations in pipette resistance. Ιn later experiments, pipettes were heat polished. This was observed at a magnification of 200x with an inverting microscope. Tips were brought to within 20-30 μm of a 25 μm Pt wire heated to a glow. Pipette tips were polished when the sides were seen to collapse slightly. Pipettes were backfilled with solution using a fine gauge hypodermic needle. Pipettes with resistances of 1-5 $M\Omega$ were used. They were only used

Figure 5. Schematic of recording configurations with a suction pipette. The uppermost frame represents a suction pipette in simple mechanical contact with a cell membrane. Upon slight suction, the seal between the membrane and the pipette increases in resistance into the giga-ohm range. Further suction disrupts the patch of membrane spanning the pipette tip. Membrane currents can be recorded under voltage clamp from whole cells after disruption of the membrane patch provided that cells of sufficiently small diameter are used. The pipette can also be withdrawn from the cell to form an outside-out patch of membrane (cyto-plasmic side of membrane is facing pipette solution).



once since debris on the tips prevented formation of giga-ohm seals. Pipettes were mounted in a microelectrode-holder with a suction port (EH-900R, W.P. Instruments, New Haven, CT). A two foot length of polyethylene (PE) tubing (I.D. 0.032") was connected to the suction port and a three-way teflon stopcock. A 1 ml gas tight syringe (Hamilton #1001, A-M Systems, Toledo, OH) and another length of PE tubing were also connected to the stopcock. The pipette holder was carefully backfilled with solution ensuring that no bubbles remained in the shank. The holder was then connected to a preamplifier (A-M Systems, Seattle, WA); the headstage of which was mounted on a micromanipulator (Leitz, Wetzlar). The micromanipulator and the inverting microscope were mounted on an anti-vibration table (Micro-g isolator, Technical Manufacturing Co., Woburn, MA).

Voltage clamp set-up: The voltage signal from the pipette was led to the input of, an electrometer, with an active bridge circuit. The membrane potential signal was compared to a command voltage using a differential amplifier. The 'error' signal was applied to the suction pipette via the current passing section of the electrometer. Rectangular pulses were generated by a WPI digitimer. Current was collected from the bath by a current voltage converter. In some experiments, a converter with a feedback resistor of 'GΩ and a bandwidth of about 4 kHz was used. In experiments, where current levels were below 1 nA, a converter with a 10 GΩ feedback register and a bandwidth of

about 2 kHz was used (built by Dr. T. Iwaszumi, University of Texas Medical Branch). The voltage clamp circuit was designed by Dr. D.C. Eaton (UTMB) (Eaton, 1972). With the system adjusted, membrane potential could be made to settle to the command voltage within 1 ms. Membrane potential and current signals were monitored by a digital oscilloscope (Nicolet, Madison, WI) and stored on magnetic diskettes. A block diagram of the potential recording and voltage clamp circuits is shown in Figure 6.

Single channel events were recorded under voltage clamp from outside-out membrane patches (Figure 5). The pipette holder was connected to the current-voltage converter (10 G Ω feedback resistor) and the reference electrode in the bath was connected to the input of the electrometer. Bath potential could then be clamped to various potentials.

Solutions

External and pipette solution are listed in Table 4. The composition of the pipette solution mimicked intracellular ionic composition, i.e. low Ca²⁺ and high K²⁺ concentrations. All solutions were made with distilled water and the pH was adjusted to 7.4. Chemicals were purchased from local suppliers if the supplier is not listed. Tetrodotoxin (TTX) and CdCl₂ were purchased from Sigma (St. Louis, MO). SITS (4-acetamino-4'-isothiocyano-2,2'-disulfonic acid stilbene) and DIDS (4,4'-diisothiocyano-2,2'-disulfonic acid stilbene) were purchased from Pierce Chemical Co.

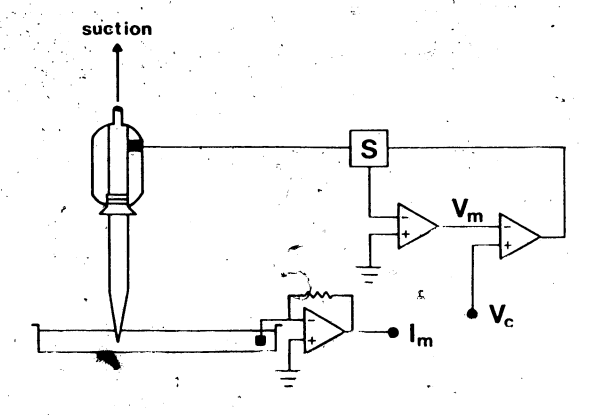


Figure 6. Experimental arrangement for voltage clamp recording. A single suction pipette is used to record voltage and pass current simultaneously. The rectangular box, denoted S, is the port on the electrometer which permits current injection. Macroscopic currents were collected from the bath by a current voltage converter (Imin pA). Vm is the membrane potential recorded by the electrometer (mV). Vc is the command voltage fed into the differential amplifier of the voltage clamp circuit (mV)

. Table 4. External and pipette solutions for whole-cell recording. Various concentrations of external Cl were made by mixing the appropriate proportions of normal and Cl Tree solutions. All values are in mM. . .

EXTERNAL

			*		EGTA (Na salt)	7	. 2	~	
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	NaCl	144	1		Nac	2	!	1,	
		Norma 1	C1 free	PIPETTE		Normal	K free	60 C1	

(Rockford, IL) and stored at -20° C. SITS and DIDS were made up as aqueous stock solutions and kept frozen in small aliquots until use. For Cl⁻ substitution experiments, NaCl or KCl were replaced by equal molar concentrations of methylsulphate (CH₃SO₄⁻) (Pfaltz & Bauer, Inc., Stamford, CT) (Hutter & Noble, 1960). CH₃SO₄⁻ has been reported to reduce fonized Ca²⁺ concentrations by about 20% as measured with Ca²⁺-sensitive microelectrodes (Kenyon & Gibbons, 1977). Therefore, the Ca²⁺ concentration was increased in CH₃SO₄⁻ solutions to keep the final concentration of ionized Ca²⁺ around 1.8 mM. Pipette solutions were filtered with a membrane filter (0.2 µm).

Prior to experiments, tissue culture dishes were rinsed several times with recording solution. Solution was aspirated with a Pasteur pipette to remove debris from the surface. Dishes were placed on the stage of an inverting microscope fitted with phase contrast optics (Olympus, Carsen, Don Mills, Ont.). In some experiments, the bath solution was changed by gravity-feed from an elevated reservoir. The volume of solution in the culture dish was reduced to less than 1 ml by inserting two Plexiglas wedges sealed to the dish with Vaseline. Complete exchange of solution could be accomplished in less than 1 minute.

Liquid junction potentials

Liquid junction potentials were measured for the suction pipettes. These potentials exist at the tip of

conventional microelectrodes and are brought about by the separation of ions with different mobilities at the interface of the pipette and bath solutions. Pipettes were immersed in the same solution as they were filled with and the potential was set to zero. Bath solution was changed to various recording solutions. No change in potential was observed for any combination of recording solutions. During this procedure, a salt bridge was employed in the bath to minimize changes in reference electrode potential.

Under all recording conditions, currents and potentials across the membrane were described with the usual convention. The external side of the membrane was taken as reference. Currents flowing from the internal to the external side were positive and displayed upwards.

All experiments were done at room temperature.

C. RESULTS

Whole-cell patch clamp technique

The whole-cell recording configuration of the single suction pipette voltage clamp (Hamill et al., 1981) was utilized to control the membrane potential of myoballs. In order to achieve adequate voltage clamp of the surface membrane potential, several conditions had to be met: 1) the seal resistance between the pipette and the cell membrane had to be high; 2) the resistance in series with the cell membrane had to be low enough to be ignored or compensated

for and 3) the intracellular space had to be isopotential.

An equivalent circuit for pipette and cell membrane is shown in Figure 7.

of giga-ohm seal: Slight positive Establishment pressure was applied to a suction pipette via a syringe and the pipette was lowered into the bath solution. The positive pressure forced solution out of the pipette and served to keep debris off the tip. For voltage clamp experiments, the clamp circuit was activated and the potential was adjusted to give zero current. This is termed zero current potential and was taken a ference potential for later measurements. A small rectangular voltage pulse was applied to the pipette and the resistance monitored continuously. Under a magnification of 200x, the tip was gently pressed against a myoball membrane and the positive pressure was relieved. Occasionally, a giga-ohm seal formed at this time as judged by a large increase in pipette input resistance to giga-ohm range (Figure 8A). Most of the time though, negative pressure had to be applied to the pipette interior by mouth to cause a giga-ohm seal to form. The development of seals occurred abruptly within seconds of application of negative pressure. After the input resistance of the pipette increased, the voltage step was increased so that the resistance of the pipette-membrane seal could be measured. Seals in the range of 1-10 $G\Omega$ were obtained routinely with pipettes that were not heat polished. Heat polished pipettes gave seal resistances of over 10 G Ω .

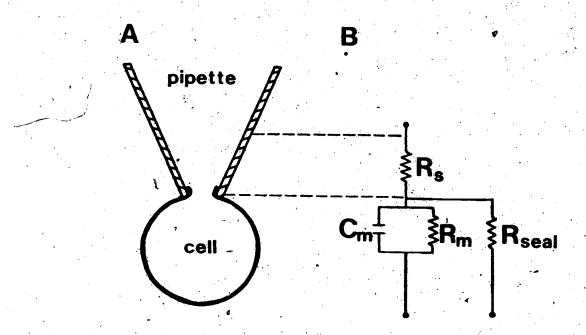
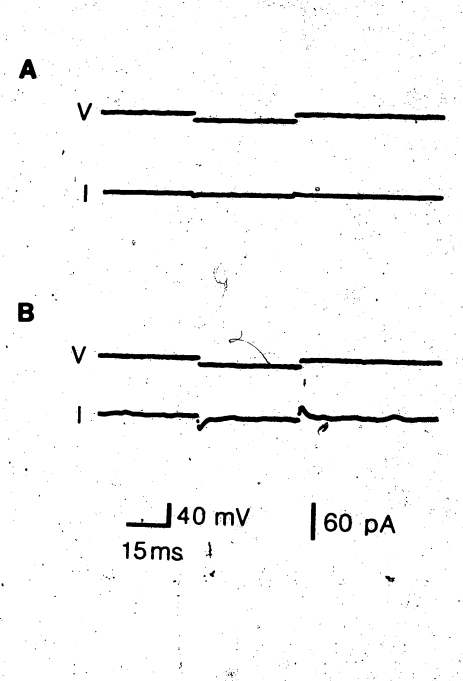


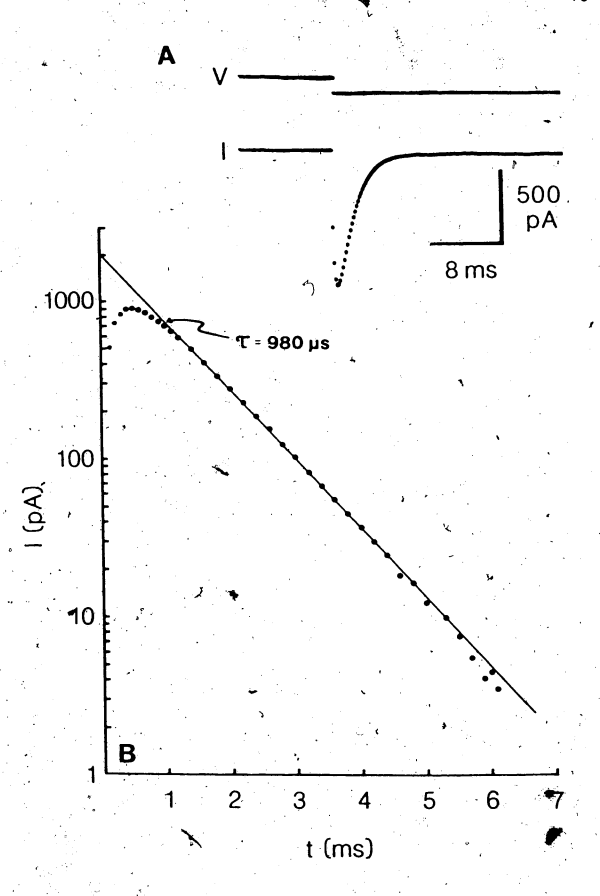
Figure 7. Equivalent circuit for pipette and cell membrane. A diagram of a cell attached to a suction pipette in the whole-cell recording configuration is shown in A and the equivalent circuit is shown in B. R. is the resistance in series with the cell membrane (M Ω). R_{seal} is the shunt resistance of the cell membrane-pipette seal (G Ω). C_m and R_m are the capacitance (μ F/cm²) and the resistance ($k\Omega$ cm²) of the cell membrane, respectively.

Figure 8. Currents recorded at different stages of whole cell recording. Establishment of a giga-ohm seal was observed as a large increase in the input resistance of the pipette to the giga-ohm range (A). Response to a hyperpolarizing voltage step indicated a seal resistance of about 3 GO. After applying further suction, rupture of the membrane patch spanning the pipette tip was indicated as a slight decrease in input resistance and an increase in (B). The capacitative current capacitative currentincreased because the capacitance of the cell now had to be Input resistance of the myoball was 1.97 G Ω . The charged. myoball had a diameter of 16 µm which gave a surface area of 804 µm assuming a spherical shape. Multiplying the input resistance by the surface area gave a unit membrane resistance of 1.5.8 k\Ocm2. The background noise was larger for the whole-cell recording configuration (B) than before rupture of the membrane patch (A) because of the additional capacitance (cf Fenwick, Marty & Neher, 1982). Input resistance of the pipette was 5 $M\Omega$ before touching the myoball.



Measurement of series resistance: To disrupt the membrane patch spanning the pipette tip, brief pulses of negative pressure were applied by mouth. Small voltage jumps showed that the disruption of the initial patch was accompanied by an increase in the input capacitance as the cell membrane now had to be charged (Figure 8B). Cell capacitance, C, could be simply degived from the total amount of charge displaced, AQ, during a step command of amplitude, $\triangle V$, according to $C = \triangle Q/\triangle V$, after subtraction of pipette capacitance. Pipette and holder capacitance were determined before disruption of the membrane patch and typically ranged from 3-10 pF. For the data shown in Figure 9, cell capacitance was calculated to be 56 pF. For 8 myoballs, the value was 58 ± 28 pF (% \pm S.D.). Assuming a spherical shape, a unit capacitance of 7 µF/cm2 was obtained for the myoball in Figure 9. For 8 myoballs, the value was $6.4 \pm 3.0 \ \mu F/cm^2 \ (x \pm S.D.)$. Unit capacitance of cultured chick myotubes has been estimated at 3.9 $\mu F/cm^2$ (Fischbach, Nameroff & Nelson, 1971). The unit capacitance of a cell membrane is about 1 μ F/cm² (Hodgkin, Huxley & Katz_k 1952). Thus, for cultured chick muscle, there appears to be a small contribution from transverse tubular membranes or from infoldings in the surface membrane. The input resistance of the myoball (Figure 9) was meas d from the steady-state current level during the voltage response and was 1.56 $G\Omega$. This gave a membrane resistance of 12.5 $k\Omega cm^2$. For 12 myoballs, membrane resistance was calculated to be

Figure 9. Capacitative current under voltage clamp. Current response to a hyperpolarizing voltage step of 44,6 mV is shown in the inset (A). Semilogrithmic plot of current as a function of time gave a single exponential fit with a time constant of 980 μ s (B). From the integral of the capacitative current recorded before the membrane patch was broken (data not shown), the capacitance of the pipette.and pipette holder was determined to be 4 pF. From the integral of the capacitative current shown in A, a total capacitance of 60 pF was obtained. Subtracting pipette capacitance from this value, a cell capacitance of 56 pF was obtained. The time constant of the declining phase of the capacitative current together with the value for cell capacitance gave a value of 17 MΩ for the series resistance. The input tance of the pipette before touching the cell was 5 MQ. The DC currrent response (A) was 28 pA giving a value of 1.6 G Ω for the input resistance of the myoball. Diameter of the 'myoball was 16 μ m. Unit capacitance was calculated to be 7 $\mu F/cm^2$. Membrane resistance was 12.5 $k\Omega cm^2$.



9.9 \pm 4.3 k Ω cm² while the input resistance was 1.2 \pm 0.6 G Ω (R \pm S.D.).

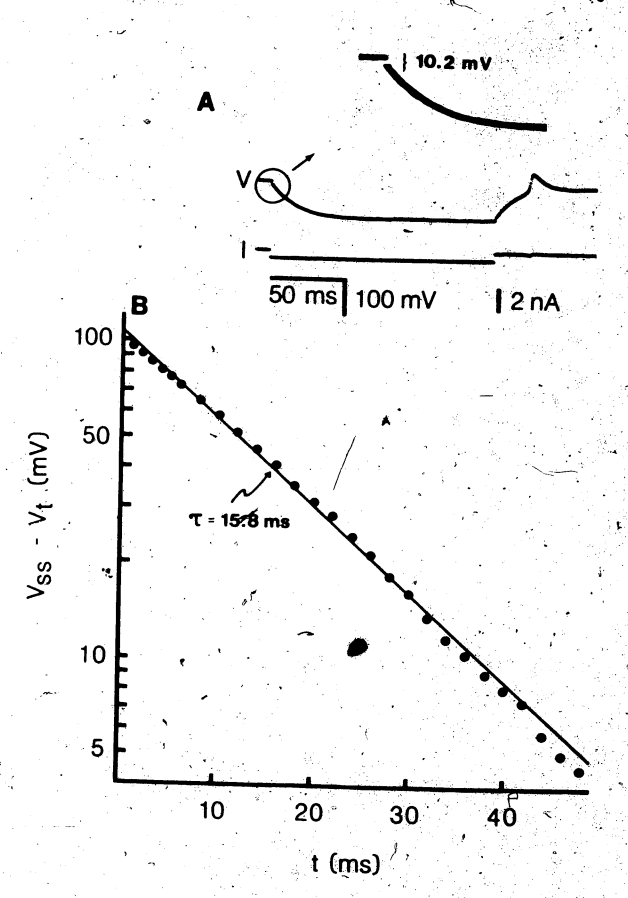
The value of the series resistance (R.) could be determined from the time constant of the decline of the capacitative current according to R. = τ/C (Marty & Neher, 1983). In Figure 9B, the capacitative current decayed exponentially with a time constant of 980 μ s. R. was calculated to be 17 M Ω . The input resistance of the pipette was 5 M Ω before touching the cell. In the best cases, R. was 3-5 times the input resistance of the pipette.

An alternative method for estimating the value of R, was to apply a relatively large hyperpolarizing current pulse to the pipette (current-clamp) after the membrane patch had been broken. The voltage drop across R, was manifested as an instantaneous voltage drop at the break of the pulse (Figure 10). For the myoball shown in Figure 10, the voltage drop was 10.2 mV giving a value of 16 M Ω for R,.

The ability thouse a single pipette to measure voltage and pass current simurtaneously depends on the magnitudes of R. and of the membrane currents. If the product of R. and current is large, then the true potential across the membrane will deviate from the command voltage by that amount. For this reason, it was important to minimize the value of R. The following precautions were taken: 1) use of low resistance pipettes and 2) selection of small diameter myoballs. Occasionally, capacitative transients were long

Figure 10. Voltage response under current clamp. A rectangular hyperpolarizing current pulse was injected and the voltage response recorded (A). Gaps in the voltage response immediately following the turn-on and turn-off of the current step (see enlargement) were the result of voltage drops across the resistance in series (R.) with the myoball membrane. The instantaneous voltage drop was 10.2 mV for a 620 pA current step giving a R, value of 16 M Ω . The input resistance of the pipette was 5 $M\Omega$. The imput resistance of the myoball was 155 $M\Omega$ and the diameter was membrane resistance was calculated to be 3.7 $k\Omega cm^2$. Semilogrithmic plot of voltage as a function of time (B) gave a single exponential. This indicated that the intracellular. space was isopotential. The measured time constant (τ) was 15.8 ms. Combined with the value for membrane resistance, the unit capacitance was calculated to be 4.3 $\mu F/cm^2$ total capacitance of 103 pF.





indicating a large R. Repeated applications of pulses of suction usually resulted in a sudden acceleration of the capacitative transient. This suggested that the slow time course was the result of incomplete rupture of the membrane patch or of clogging of the pipette tip with intracellular material. Most of the time, this procedure brought about a stable change in the transient. Data was only accepted for analysis is the product of R. and maximal current was less than 3 mV. No R. compensation (Hodgkin & Huxley, 1952a) was employed as this restriction could generally be met.

Spatial control of voltage: Another important consideration was spatial control of voltage. Because of the long, tubular morphology of muscle cells (myotubes) grown in fissue culture, the cable properties of these cells prevent adequate nontrol of membrane potential under voltage clamp (Fuhuda et al., 1976b). To avoid this problem, the shape of the muscles cells was altered by including colchicine in the time another me is (Fukuda et al., 1976a). When grown in the time another of colchicine, the macle ells assumed a coulty charter (Flate al.) The effect of colchicine and a problem of the colchicine of the colchicine

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(Figure 10B). As well, the capacitative current recorded under voltage clamp had an exponential decline (Figure 9B). Spatial control of voltage was also reflected in current voltage plots of time-dependent currents (cf weck et al., 1975). Adequate spatial control was reflected in current voltage plots if the curve was nearly symmetrical around the inward current. Whereas peak current levels were attained within a few millivolts of threshold if voltage was poor (see Figure 39). This is because the control voltage clamp cannot oppose the depolarization caused by the opening of channels. In general, current-voltage plots of inward currents were symmetrical. Poor voltage control was also apparent from the membrane voltage traces as deviations from the rectangular shape. This was generally not seen except when membrane currents were very large (several $n\lambda$).

Exchange of pipette and intracellular solutions: The wholencell recording technique offered the opportunity to alter intracellular ion commosition and to apply drunk to the cytoplasmic side of the membrane. There was every evidence that, provided the capacitative transient was fast, exchange on urred between pipette and mychall colutions within about the minute. The mychalle had no tward currents (Figure 11) which quickly distributed if the ripette was filled with K free solution (Ch. replacement) containing totraetby armonic (TEA). For the TOP and the are known to

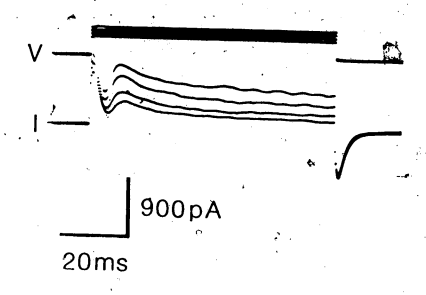


Figure 11. Outward membrane currents. Depolarizing voltage steps were applied from a holding potential of -60 mV. Four traces are superimposed. The currents appeared to exhibit voltage dependent inactivation. Repetitive voltage steps (every 200 ms) reduced the outward current level. After a several second rest, the current level returned to the initial level (data not shown). External solution was normal containing 10.7 M 100 mM cd. Pipette solution was

(Hille, 1970) and the outward currents were most likely carried by K. Also, as will be shown later embrane Cl-currents could be made to reverse at different potentials by altering the intracellular concentration of Cl-.

'It was my experience that the use of suction pipettes to penetrate a cell inflicted much less damage than conventional microelectrode impalement. Whole-cell recording could be performed for periods up to one hour without visual or electrical signs of deterioration.

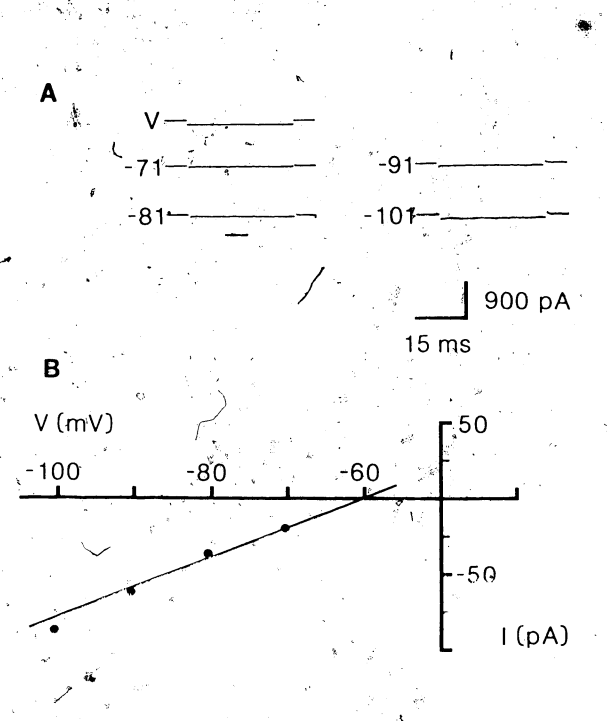
In summary, the use of single suction pipettes for whole-cell voltage clamp had the following characteristics:

1) adequate control of membrane potential; 2) opportunity to alter intracellular ion concentrations and to apply drugs to the internal surface of the membrane and 3) minimal damage to the cells. In addition, the membrane-pipette seal was mechanically stable. This allowed the formation of cutside-out patches for the recording of single channel currents (see Discussion).

Membrane currents under woltage clamp

Leakage currents and must be subtracted from time-dependent currents. Leakage conductance was assumed to be linearly related to potential over the range of potentials at which time-dependent currents were studied. Leakage conductance was determined by applying small, hyper-

Figure 12. Determination of leakage conductance. Records of voltage and current during hyperpolarizing voltage steps from a holding potential of -60 mV (A). Potential to which the membrane potential was stepped is indicated beside each current trace. Current magnitude was plotted as a function of test potential (B). The slope of the line gave an input resistance of 0.55 G Ω . Myoball diameter was 25 μ m. Value calculated for membrane resistance was 10.8 k Ω cm². The seal resistance was 2 G Ω .



was constructed and conductance was determined from the slope of the line. This value was used to calculate leakage currents for depolarizing voltage steps.

Voltage- and time-dependent currents: Under voltage clamp conditions and with the pipette and bath solutions approximating normal somposition. ionic responses to depolarizing voltage steps were recorded (Figure 13). There was a fast inward current which was blocked by 10-7 M TTX and was presumably carried by Na+ (Narahashi et al., 1964). At more depolarized potentials, outward currents developed. They were probably carried by K since the outward currents were absent when the pipette solution contained (Stanfield, 1970b; Armstrong, 1975). The slaw inward current and the slow inward current flowing during repolarization to the holding potential were identified as Cl currents. They are described in more detail in the following sections. Ca 2+ \$ currents have also been reported in these cells (Fukuda et al., 1976b), but they were probably masked by the other currents. Under current clamp, a long duration action potential (Figure 14A) and a short, fast spike (not shown) 'observed.

Voltage- and time-dependent chloride currents: To isolate the current flowing through the voltage-dependent Cl conductance, currents flowing through the other permeability pathways were minimized by including channel blockers in the solutions and by substituting impermeant ions for

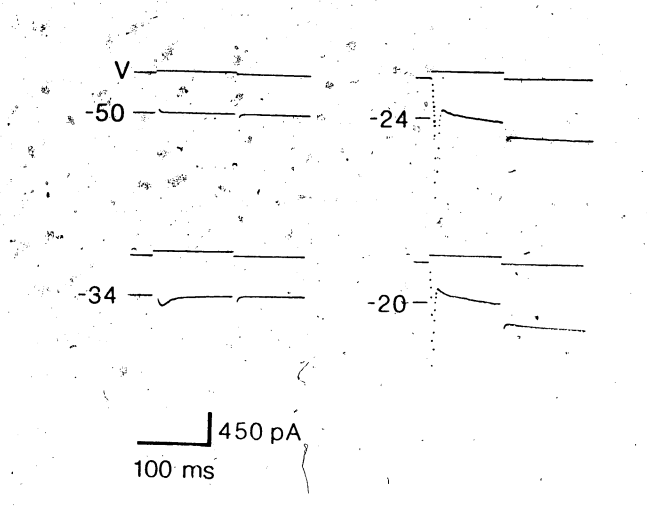
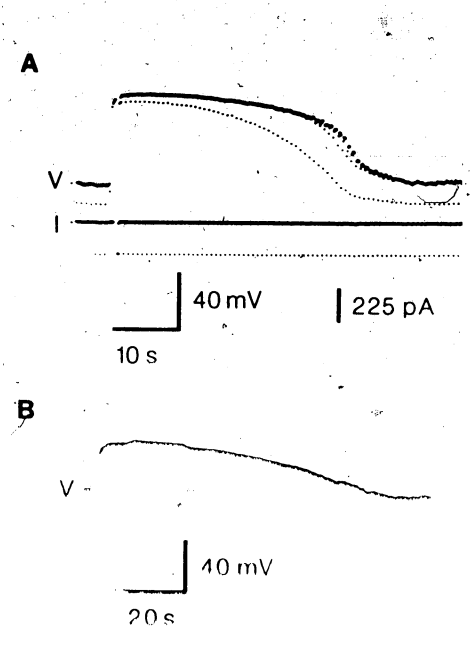


Figure: 13. Membrane currents under voltage clamp. Depolar izing voltage steps were applied from a holding potential of -60 mV. The potential to which the membrane potential was stepped is indicated beside each current trace. External and pipette solutions were normal that two current traces recorded.

Figure 14. Chloride action potentials. Under current clamp, a long duration action potential was elicited (A). Small hyperpolarizing current pulses were given continuously to show the change in membrane resistance during the action potential. Resting input resistance was 0.6 G Ω . At the top of the plateau of the action potential, input resistance dropped to 0.2 G Ω . During the repolarization phase of the action potential, input resistance increased to 1.3 $G\Omega$. Resting potential was -70 mV. Duration of action potential was 38 s and height was 60 mV. External and pipette solutions were normal. Part B shows an action potential recorded in the presence of 20 mM TEA intracellularly. The duration was 85 s and the height was 39 mV. Resting potential was -63 mV and the potential at the top of the action potential was -28 mV. Reversal potential for Cl given by the Nernst equation was 23 mV. External solution was normal containing 10 7 H TTX and 1.0 mM Cd2. Pipette molution was 50 Cl .

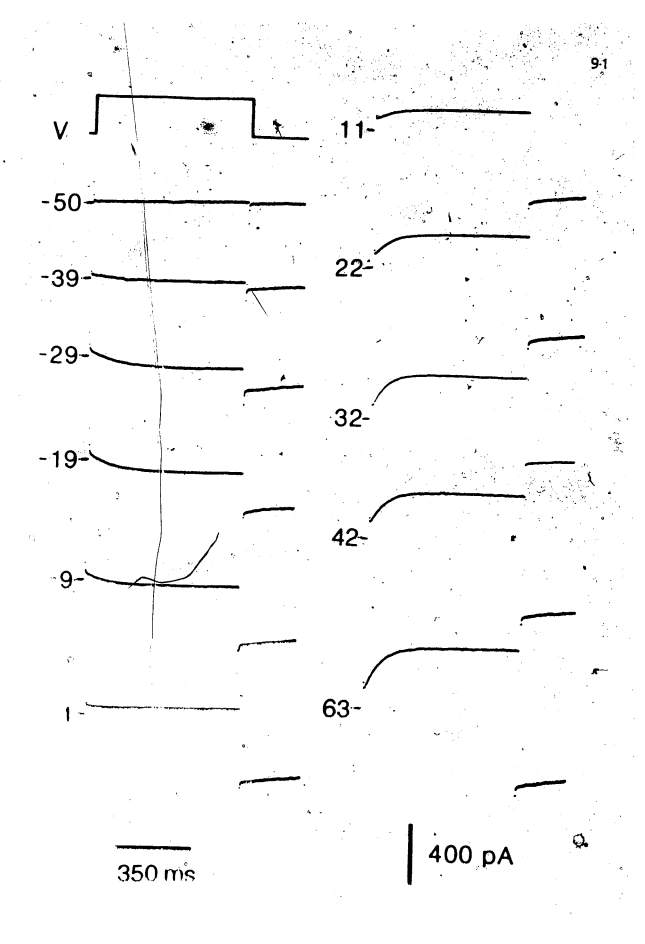


permeant ones. Na* currents were blocked by including 10-7 M TTX in the bath solution (Narahashi et al., 1964) and Ca²* currents were blocked by 1.0 mM Cd²* in the bath (Hagiwara & Byerly, 1981). K* currents were eliminated by using K*-free pipette solution with Cs* as the major cation and by including 20 mM TEA (Stanfield, 1970b; Armstrong, 1975).

Figure 15 shows records of currents elicited with ms depolarizing sweltage steps from a holding potential of -60 mV. As the potential was stepped further away from the holding potential, an inward current developed slowly and did not decline during maintained depolarization. As the potential was stepped to more positive values, the inward current rose more quickly and became larger. Near 0 mV, the current trace was flat, i.e. that was the reversal potential or the currents as no net current was flowing. This also showed that the other currents were successfully blocked. Above 0 mV currents were outward and rose more quickly than less positive potentials. Following the depolarizing steps, inward currents were recorded which did not decline for several seconds (only partially shown in Figure 15). Under current clamp, an extremely long duration action potential was recorded (Figure 14B). Cl action potentials recorded under 'normal' ionic conditions (Figure 14A) had much shorter durations. Consequently, outward K* currents must play a role in repolarizing the membrane potential.

The currents were corrected for leakage currents and steady state current levels were plotted against

Figure 15. Membrane chloride currents. Membrane currents were recorded under voltage clamp during step depolarizations (V) from a holding potential of -60 mV. Voltage steps were to the potential listed beside each current trace. Currents through Na*, and K* and Ca²* channels wre minimized by including 10^{-7} M TTX and 1.0 mM Cd²* in the external solution and by having 20 mM TEA in a K*-free solution inside the pipette. Seal resistance was 2 G Ω_{\bullet} Pipette input resistance was 2 M Ω_{\circ} R, was 6 M Ω_{\circ} Diameter of the myoball was 16 μ m. Membrane resistance was calculated to be 8 k Ω cm². External solution was no mal plus the additions mentioned above (149 mM Cl) Firette solution was K*-free (155 mM Cl) (2000)

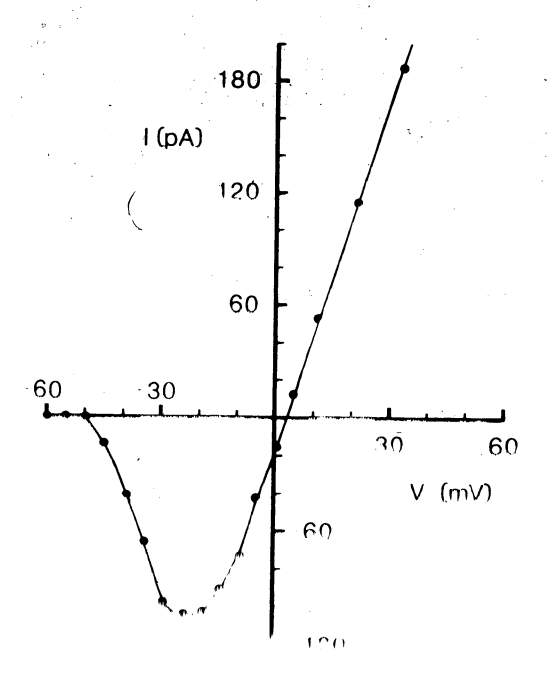


depolarizing test potential (Figure 16). Currents were detectable above -45 mV and were initially inward. Peak inward current occurred at about -25 mV and then became less inward and reversed polarity above 0 mV. Measured reversal potential was 3 mV. Peak current density was 12 μ A/cm². Peak current densities ranged from 3 to 30 with a mean of 17 μ A/cm² for 12 myoballs.

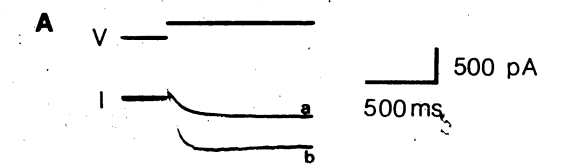
In two experiments, external () concentration was changed during the recording. When the external () concentration was reduced, inward concentration became larger indicating participation of 1 in the currents (Figure 17A).

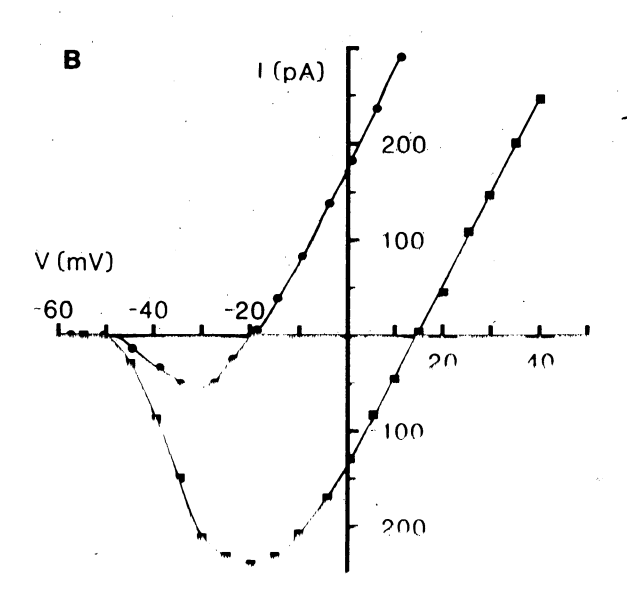
To demonstrate quantitatively that the slow membrane currents were carried by Cl., reversal potentials, for steady-state currents were measured in a mariety of internal and external Cl concentrations. Care was taken to select pipettes with wide tips and mydealls with small diameters so as to achieve the best mossible outrate intracellular ion concentrations of was r placed with the impermeant ton, methyla libáte (Kutting Michle, 1960) ''qure 178 abowa ibe results for two liffe it sob le for the is re orded in TO mM viernal and the met ate call of the measured remembed priontial was if mo, the opullibrium potential calculated from the Naruet on out of 18 mm, may per the myoball in 140 mm external and the mm internal of the mean red reversal resemble as 1. i . . . eg:

Figure 16. Current voltage plot of chloride currents. Magnitudes of steady-state currents were corrected for leakage currents and plotted as a function of depolarizing test potential Measured reversal potential was 3 mV. Peak current density was $12~\mu\text{A/cm}^2$ (measured at -25 mV). Data



Effects of changing external chloride concen-Figure 17. trations on membrane currents. Currents recorded during depolarizing voltage steps to -19 mV from a holding potential of 60 mV (A). External Cl concentration was reduced from 149 mM (current trace a) to 3 mM (current trace h) while continuously recording from the myoball. C1replaced with methylsulphate. was Current-voltage plots for currents recorded from different myoballs in different Cl concentrations te shown in B. The curve drawn through the circles is data from recordings done in 60 mM intracellular Cl and 149 mM external Cl., Measured reversal potential was -19 mV. The curve drawn through the squares is data from recordings done in 155 mM intracellular Cl and 76 mM external Measured reversal rotential was 15 mV. Holding potential was commy for both. External solution contained 10.7 M TTX and the errod's tip the colution contained 60 mM TEA.





The graph in Figure 18 shows the cumulative results from 21 myoballs. The reversal potential for the steady-state current was plotted as a function of Cl-concentrations. The regression line through the data points has a slope of 53 mV per decade which is close to the 58 mV per decade predicted by the Nernst equation. The slope of the line probably would have been closer to the predicted value if better control of intracellular ion concentrations was possible. This data gave very strong evidence that the slow currents were carried by Cl-.

Block by SITS, DIDS, and SCN; Stilbene derivatives have been shown to block Cl permeability in a wide variety of cell types: red blood cells (Knauf & Rothstein, 1971; Cabantchik & Rothstein, 1974); epithelial cells (Ehrenspeck & Brodsky, 1976) and invertebrate (Russell & Brodwick, 1979) and vertebrate (Vaughan & Fong, 1978) skeletal muscle. It has been suggested that SITS and DIDS act at the Cl binding site (Shami, Rothstein & Knauf, 1978). It was found that SITS (1 mm) blocked the currents in a reversible manner (Figure 19). DIDS (10 μM) also blocked the currents, but the block could not be reversed by repeated washing. irreversibility of the block by DIDS has also been observed in other preparations (Cabantchik & Rothstein, 1974). SCN-(10 mM) was found to block the currents in a reversible manner as has been reported for the Cl channel in electroplax membranes (White & Miller, 1981).

Figure 18. Reversal potentials as a function of chloride concentrations. Reversal potentials for atteady-state currents were measured in various Cl concentrations as demonstrated in Figure 17. Data from 21 myoballs. Values are means ± S.D. The straight line was drawn according to the Nernst equation:

$$E_{Cq} = \frac{RT}{zF} ln \frac{(Cl)}{(Cl)}$$

 E_{Cl} is the equilibrium potential for Cl^- (mV); (Cl)_o and (Cl)_i are external and internal Cl^- concentrations (mM), respectively. R, T, z and F have their usual meanings.

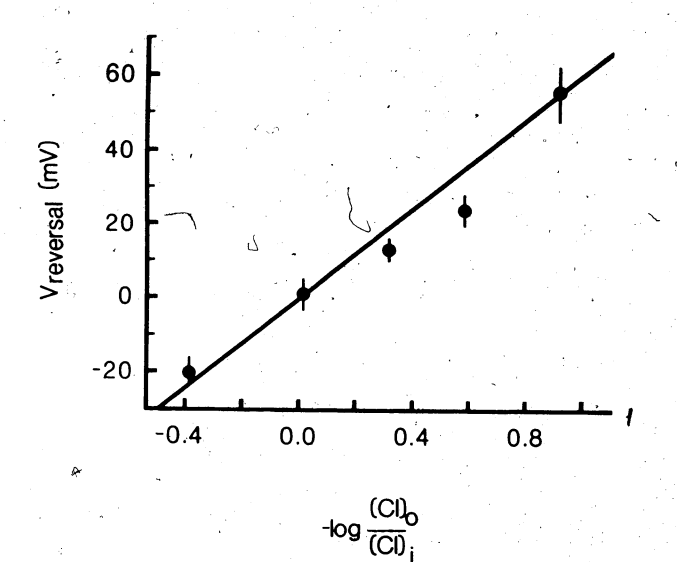
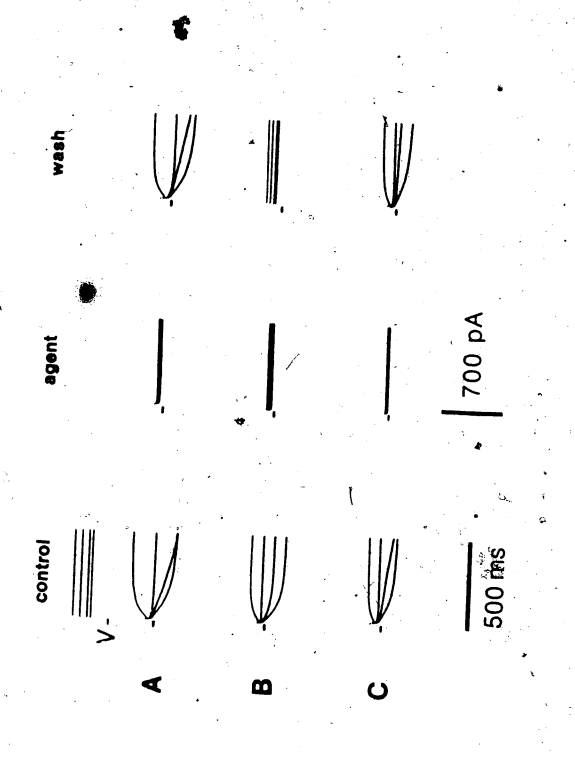


Figure 19. Effects of SITS, DIDS and SCN on membrane currents. Currents were recorded under voltage clamp. A control series was taken in normal external solution containing 10-7 M TTX and 1.0 mM Cd²⁺. Pipette solution was K⁺-free. Bath solution was then changed to one containing the different agents while continuously recording from the same myoball. Four current traces are superimposed in each frame. The voltage step protocol is denoted V. Currents were reversibly blocked by 1 mM SITS (row A). Block by 10 μ M DIDS was apparently irreversible as extensive washing with normal external solution did not reverse the effect (row B). Block by 10 mM SCN was reversible (row C). Holding potentials were -60 mV in B and -80 mV in A and C.



C1 currents lack an inactivation mechanism: Since the currents did not decline during maintained depolarization (see Figure 15), this indicated that the conductance lacked an inactivation mechanism. To test whether there was any steady-state voltage dependent inactivation operating at negative potentials, the holding potential was varied and depolarizing voltage steps to the same potential were applied (Figure 20). It can be seen that the magnitude of the inward currents elicited by the test pulses did not change. Therefore, over the range of potentials tested, there was no evidence of voltage-dependent inactivation.

Tail currents: Following depolarizing voltage steps that activated Cl currents, currents became inward and decayed slowly and completely (Figure 21). Currents that flow during repolarization are termed tail currents. They were examined with a two pulse protocol. First, a large depolarizing voltage step was applied in order to open as many channels as possible. Then the potential was stepped back to various levels (Figure 22A). Magnitudes of the tail currents were measured following the break the depolarizing They were plotted as a function of repolarization potential in order , to construct an 'instantaneous' current-voltage plot (Figure 22B). Data points could be fitted a straight line with an extrapolated reversal potential of I mV. This value was very close to the calculated Cl equilibrium potential of 1 mV and indicated that the tail currents were carried by Cl.. The straight line fit to

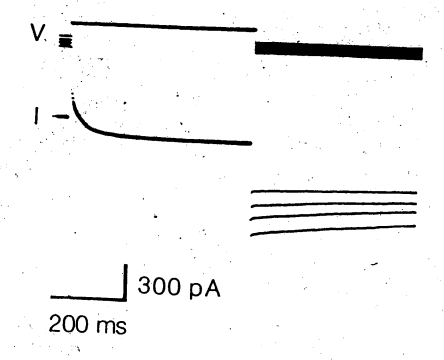


Figure 20. Lack of voltage-dependent inactivation. Cl-currents were recorded under voltage clamp. The holding potential was varied, -60, -70, -80 and -89 mV, and a depolarizing voltage step to -23 mV was applied. Four traces are superimposed. Note that the steady-state current level was identical in all four runs. External solution was normal containing 10⁻⁷ M TTX and 1.0 mM Cd²⁺. Pipette solution was K*-free.

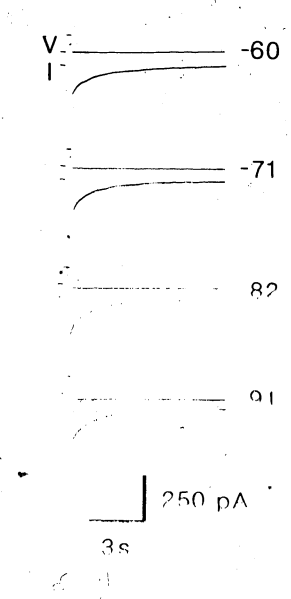
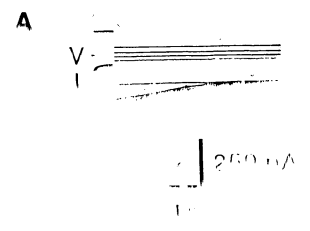
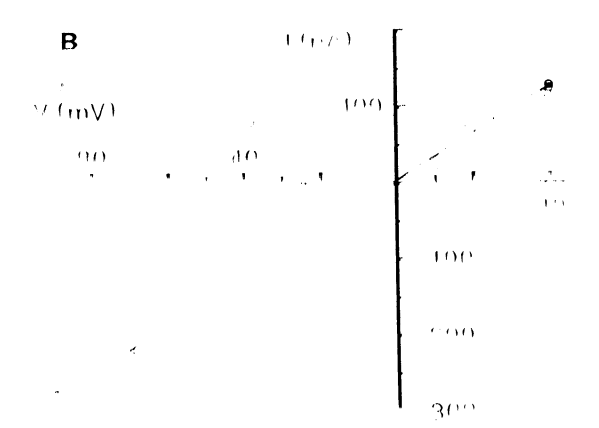


Figure 21. Tail currents. Currents flowing during repolarization to carious potentials following a depolarizing step to 38 mV for 300 ms are shown. Potential during repolarization: is listed beside each trace. Holding potential was 60 mV. External solution was normal containing 10 M TTX and 1.0 mM Cd Tripette solution was Kinfron. Seal resistance was 4 CO. No. 1992 2 MM torus resistance was 4 CO. No. 1992 2 MM torus resistance of the pipette was 2 MM

Figure 22. Instantaneous current-voltage plot. Four superimposed traces show the protocol for collecting data for instantaneous current-voltage plots (A). Following a depolarizing voltage step to 40 mV, the potential was stepped to various potentials. Tail current magnitudes were measured following the break of the depolarizing pulse and corrected for leakage conductance. Values were plotted against repolarization potential (B). Data from are represented by filled circles and from steady state currents by circles. Data points fell on a straight line with a slope of 3.2 nS. Diameter of myoball was 13.6 μm which gave a conductance of 0.55 mS/cm². Measured reversal potential was -1 mV. Holding potential mV. External solution was normal containing 6.0 M TTX an' 1.0 mm rd2'. Pipette solution was K'-free. was 8 MO. Rin of pipette Scal * 7101



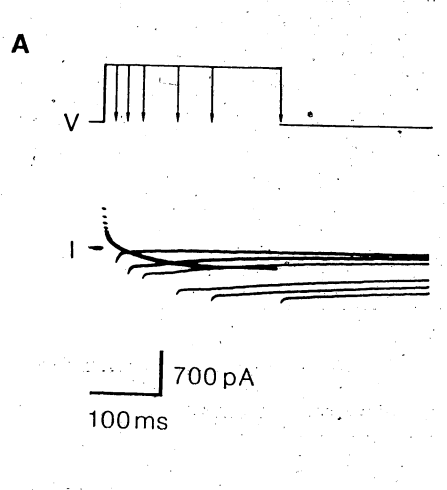


the data also indicated that the open channels behaved ohmically. The slope of the line gave a conductance value of 3.1 nS. The mean value for 10 myoballs was 8.3 ± 3.9 nS (9.4 + 5.0.) Calculated per unit area the mean conductance was 0.63 ± 0.3 mS/cm².

Tail currents were also examined by varying the duration of the depolarizing test pulse (Figure 23). Membrane conductance was calculated from the steady-state and the tail currents at the end of the depolarizing test pulse. The values were plotted as a function of time and it can be seen that the time course of the conductance changes calculated from both steady state and tail currents was similar. This result supported the idea that the tail currents were floring through the conductance mechanism activated during the depolarizing pulse.

Cl conductance as a function of voltage: Since the open channels were found to behave obmically, the following relaticumhir could be applied: $I = G(" + V_{rev})$, r is current $(pA)_{+}$ G is conductance $(mS)_{+}$ V is voltage at which current was monoured (mt) and t is the reversal potential (mv) relation, conductione was galculated this normáli em with respect to the maximum value for myohall (Figure 24). The resulting curve reflects the fraction of channels open at a given membrane potential. The relationship, between conductance and voltage was sigmoid. It rose shamply between 40 and 0 mV and levelted off remit or a toutified whe mean reduction maximal Cl

Figure 23. Envelope of tails. Currents recorded in response to depolarizing steps to -29 mV from a holding potential of -60 mV are superimposed (A). The duration of the depolarizing pulse was lengthened as indicated in diagram (V). Pulse durations shown are: 12, 30, 50, 100, and 250 ms. Conductance was calculated termination of the depolarizing pulse from both steady-state currents (circles) and from the tail currents (triangles) (B). The values are plotted as a function of time. Note that the time course of the conductance changes from either the steady-state or the tail calculated currents followed a similar time course. Reversal potential External solution was normal containing mV. 10 7 M TTX and 1.0 mM Cd2. Pipette solution was Ki-free. Seal resistance was 5 GO. R. was 6 MO. R., of pipette was 1 MΩ.



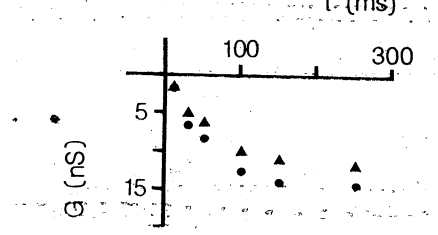
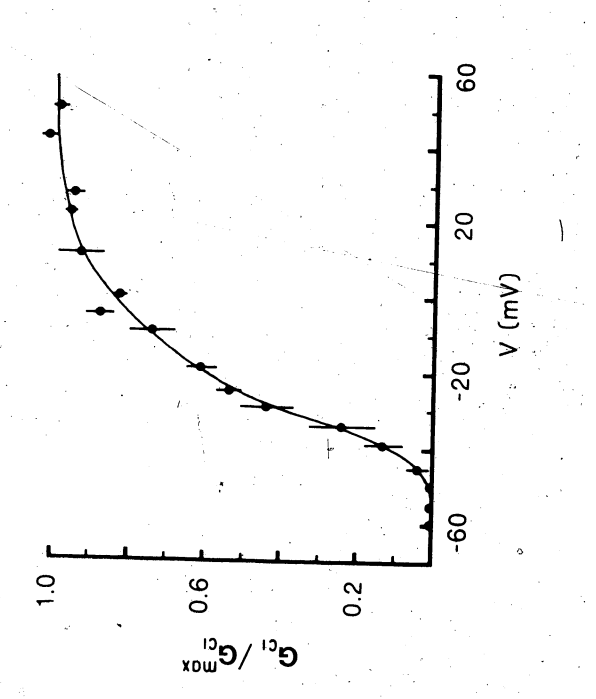


Figure 24. Conductance as a function of voltage. Conductance was calculated according to: $I = G(V - V_{rev})$. I is current (pA); G is conductance (nS); V is membrane potential (mV); V_{rev} is measured reversal potential (mV). Conductance values (G_{Cl}) were normalized with respect to the maximal conductance (G_{Cl}^{max}) for each myoball. Data from 5 myoballs. Values are means \pm S.E. Smooth curve drawn by eye.



conductance was 1.03 \pm 0.7 mS/cm² (R \pm S.D.; n = 13).

The relationship between Cl conductance and membrane potential could also be seen in the current-voltage plots, if the magnitudes of the tail currents were measured for each of the depolarizing steps (Figure 25). The resulting tail current amplitude versus voltage curve should be proportional to the fraction of open channels at each potential. The curve resembled the curve in Figure 24 in that it rose steeply from -40 to 0 mV and then started leveling off above 0 mV.

Plotting Cl conductance as a function of voltage on a semilogrithmic plot showed that the values approached a limiting value of 8 mV for an e-fold change in conductance (Figure 26). Thus, the Cl conductance exhibited high voltage sensitivity. The Na conductance of frog skeletal muscle has been reported to rise e-fold for a 3.7 mV depolarization (Ildefonse & Rougier, 1972; Hille & Campbell, 1976).

of Cl currents as a function of time gave straight line fits suggesting a monoexponential time course for current activation (Figure 27). However, it was found that the currents could not be completely described by a single exponential as the first 6-10 ms of current development did not fit. There appeared to be delays of several milliseconds in the development of the Cl currents. Unfortunately, the capacitative transients obscured most of the delays. In

Figure 25. Current-voltage plot of tail current magnitude. Steady-state currents (circles) were plotted against depolarizing test potential. Tail current magnitudes (squares) were measured immediately after the turn-off of the depolarizing test pulse and plotted against test potential. The resulting tail current amplitude curve should be proportional to the fraction of channels that were activated at the depolarizing test potential. Compare to the conductance versus voltage curve in Figure 24. Data taken from the currents shown in Figure 15.

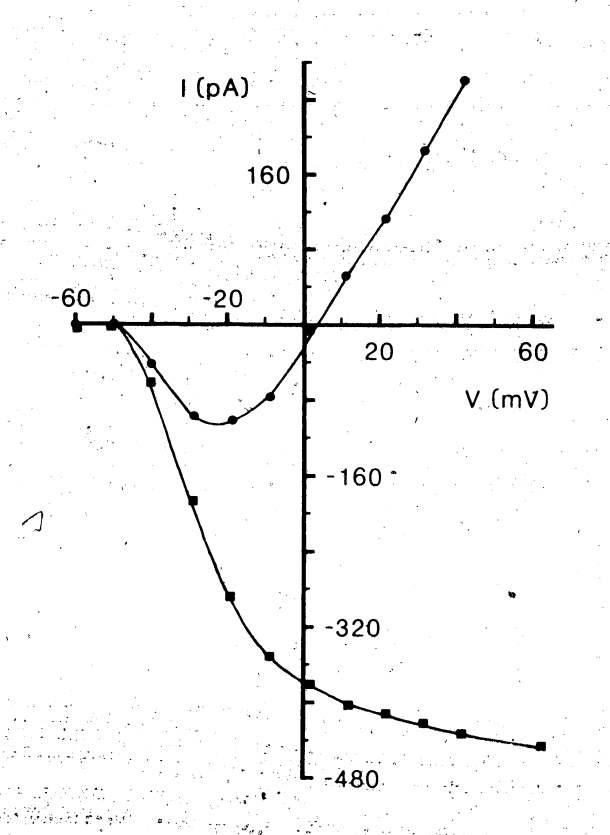


Figure 26. Semilogrithmic plot of conductance as a function of voltage. Normalized values of Cl conductance $(G_{\text{Cl}}/G_{\text{Cl}}^{\text{max}})$ were plotted against voltage on semilog paper. Data same as in Figure 24. Line through the data is a regression line fit to the conductance values for potentials more negative than -25 mV. Slope of the line gives an e-fold change in conductance for an 8 mV depolarization.

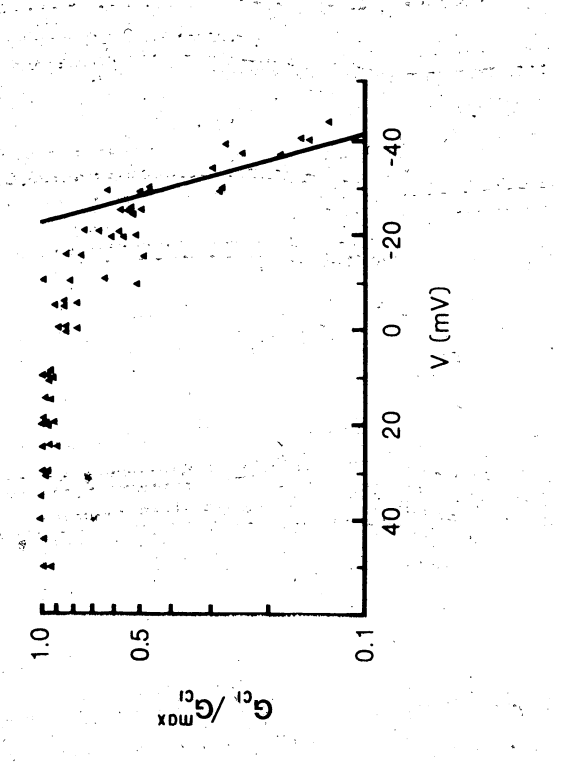
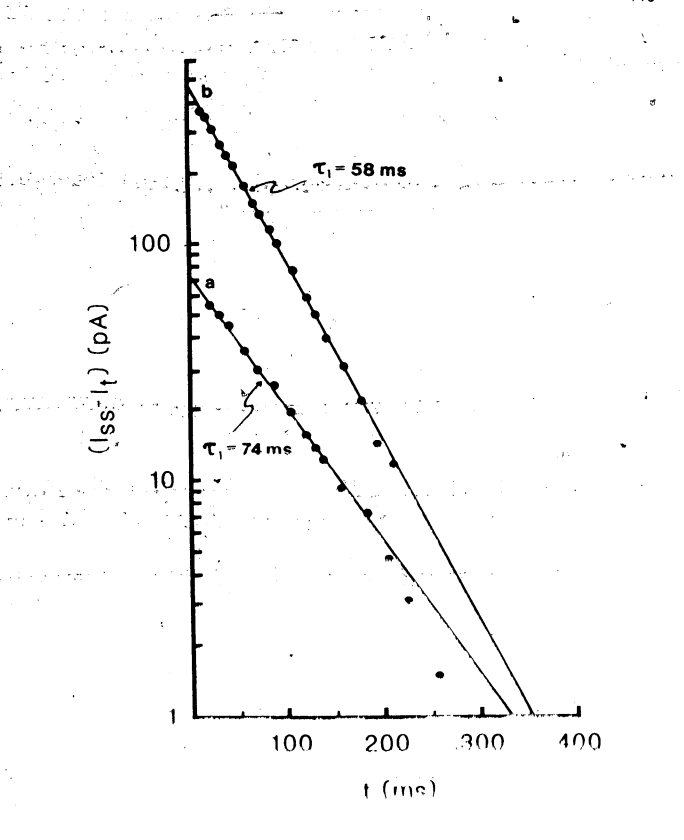


Figure 27. Semilogrithmic plot of current activation. Inward current, magnitudes for voltage steps to -9 mV (line a) and to 63 mV (line b) were plotted as a function of time. The time constant (τ_1) for the voltage step to -9 mV was 74 ms with an amplitude (A_1) of 73 pA. The time constant (τ_1) for the step to 63 mV was 58 ms and the amplitude (A_1) was 473 pA. Turn-on of the depolarizing voltage step, was set as time equal to zero. Same current traces as in Figure 15.



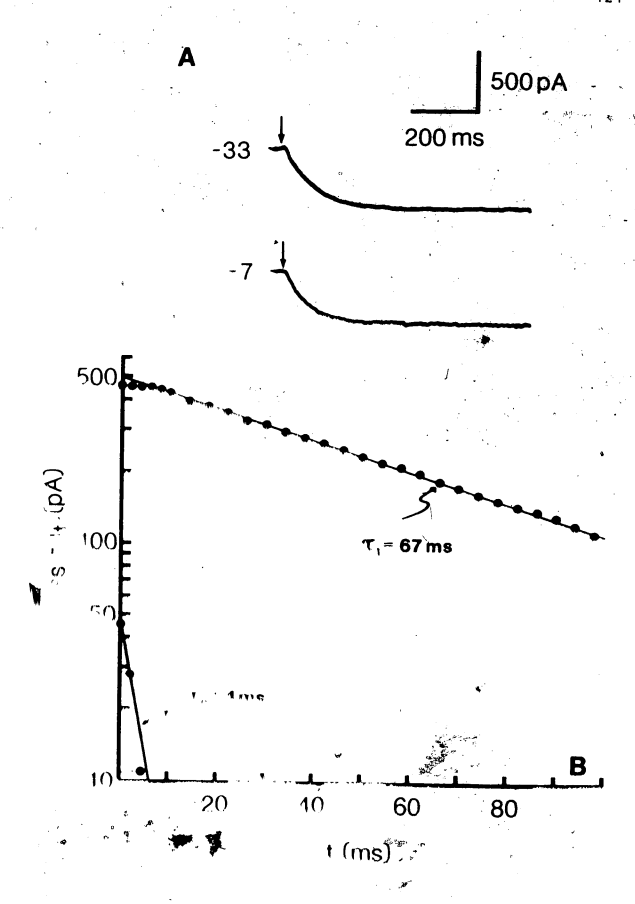
order to visualize the delays, the capacitative currents were eliminated from the current records by adding current responses to hyperpolarizing voltage steps of equal magnitude. For the traces shown in Figure 28, there was about a 6 ms delay in inward current development for a depolarizing step to -33 mV and about a 4 ms delay for a step to -7 mV. Delays were found to be slightly shorter for currents recorded at more positive potentials. Plotting current records of this type versus time on semilogrithmic plots showed that the initial slope was close to zero (Figure 28B).

To account for these observations, the time course of activation of the Cl currents was fit with the sum of two exponentials: a major component with amplitude A₁ (pA) and time constant τ , (ms) and a minor component with amplitude A₀ (pA) and time constant τ_0 (ms). The amplitude of the minor exponential component had the opposite polarity to the amplitude of the major exponential component in order to account for the delay in current activation. The sum of the two exponentials gave an initial slope close to zero. The values of A₀, A₁ and τ_1 were obtained by fitting a straight line to a semilogrithmic plot of current (see Figure 28). The was roughly estimated by fitting the model current to the actual current record. The equation used to fit the currents

 $[\]gamma_{+} = \gamma_{-}$ [Noexp(+/ τ_{0}) $\sim N_{1} \exp(-t/\tau_{1})$]

to pull to the electronic of the topological steady state,

Eigure 28. Delay of activation. Cl currents recorded in response to depolarizing voltage steps to -33 and -7 mV from a holding potetial of -70 mV are shown in Part A. Linear leakage and capacitative currents were subtracted' from the inward current recomes by adding the current responses to voltage steps opposite in polarity. Arrows indicate the turn-on of the voltage steps. There was a slight' delay in the development of the inward currents which was about 6 ms for the step to :33 mV and about 1 ms for the step to -7 mV. Semilogrithmic plot of current as a function of time for a voltage step to -18 mV was fitted with a staight line (B). The time constant for the major exponential (1.) was 67 ms and had an amplitude (1.) of 518 ph. The initial slope was close to zero. Another exponential with a time constant (70) of roughly 4 ms, and, an amplitude (A) of 46 f gave an initial clare of zero if subtracted from hold jor expression, it erns and in was a minty of early and the high



(pA), respectively. Figure 29 shows that the fit of the equation to the currents was excellent.

The time constants of the major exponential component (τ_1) were found to vary with membrane potential (Figure 30). They decreased from 150-200 ms at potentials more negative than -20 mV to about 50 ms at potentials more positive than 0 mV. The time constants (τ_0) of the minor exponential seemed to vary little with voltage, but the values were very uncertain. The presence of this component (delay) suggests that there are at least two kinetically distinct closed states.

C1 current deactivation kinetics: Tail currents had an unusually slow time course (Figure 21). Therefore, consideration had to be given to the possibility that there was a component of the tail currents that did not reflect the closing behavior of the Cl channels. There was no indication that voltage control was lost since the voltage trace was perfectly flat and the tail currents declined with exponential time courses (discussed later). Another source of artifact could have been polarization of the Aq/AgCl electrode during the prolonged current flow. However, this seemed unlikely as rolarization would be expected to distort the recording of alow and steady state events (of Katz, 1966). A more likely explanation would have been the effects of ion accumulation and/or derletion in a restricted space. The following the community of the continued to evaluate this

1 - - - - 11. 11.

Figure 29. Fit of model equation to currents. Curves modelling activation of Cl currents were drawn according to:

 $I_{t} = I_{..} - [A_{0}exp(-t/\tau_{0}) + A_{1}exp(-t/\tau_{1})].$

I, is current at time t (pA); I. is steady-state current (pA); A_0 is amplitude of minor exponential (pA); τ_0 is time constant of minor exponential (ms); A_1 is amplitude of major exponential (pA); τ_1 is time constant of major time exponential (ms).

For a depolarizing voltage step to -28 mV, I.. = -464 pA; $\lambda_0 = 54 \text{ pA}$; $\tau_0 = 9 \text{ ms}$; $\lambda_1 = -518 \text{ pA}$; $\tau_1 = 128 \text{ ms}$. For a depolarizing step to -18 mV, I.. = -452 pA; $\lambda_0 = 46 \text{ pA}$; $\tau_0 = 4 \text{ ms}$; $\lambda_1 = -498 \text{ pA}$; $\tau_1 = 67 \text{ ms}$. Turn-on of voltage step indicated by vertical bars. Data points from steps to -28 and -18 mV from a holding potential of -70 mV were superimposed on the model curves. Linear leakage and capacitative currents were eliminated by adding current responses to voltage steps of opposite polarity. Currents were from same myoball as in Figure 28.



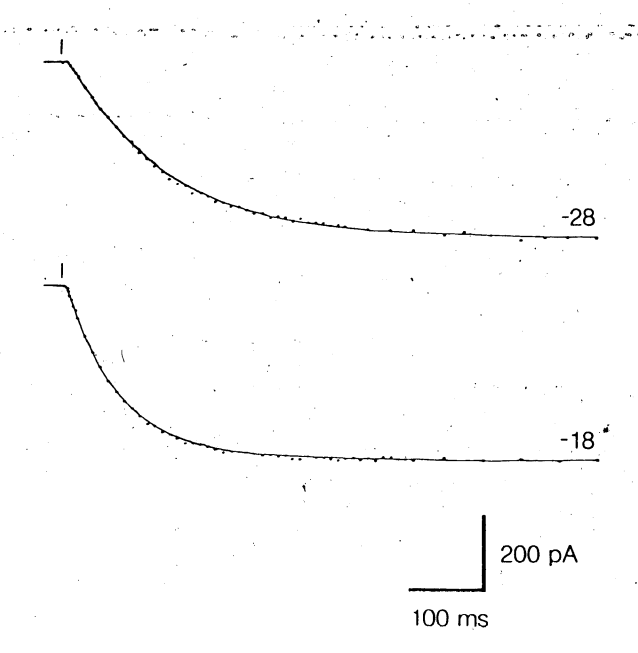
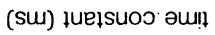
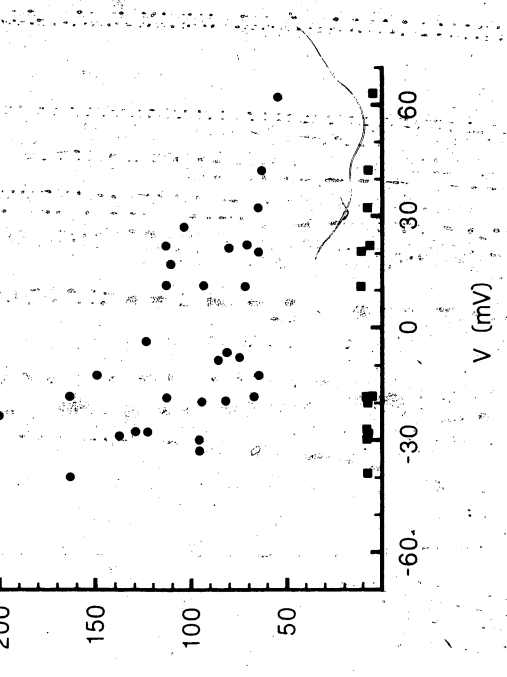


Figure 30. Time constants of activation as a function of voltage. Time constants for the activation of Cl^- currents were obtained as shown in Figures 27, 28 and 29. Circles represent time constants for the major exponential component (τ_1) . Data from 5 myoballs. Squares represent time constants for the minor exponential component (τ_0) . Data from 3 myoballs.

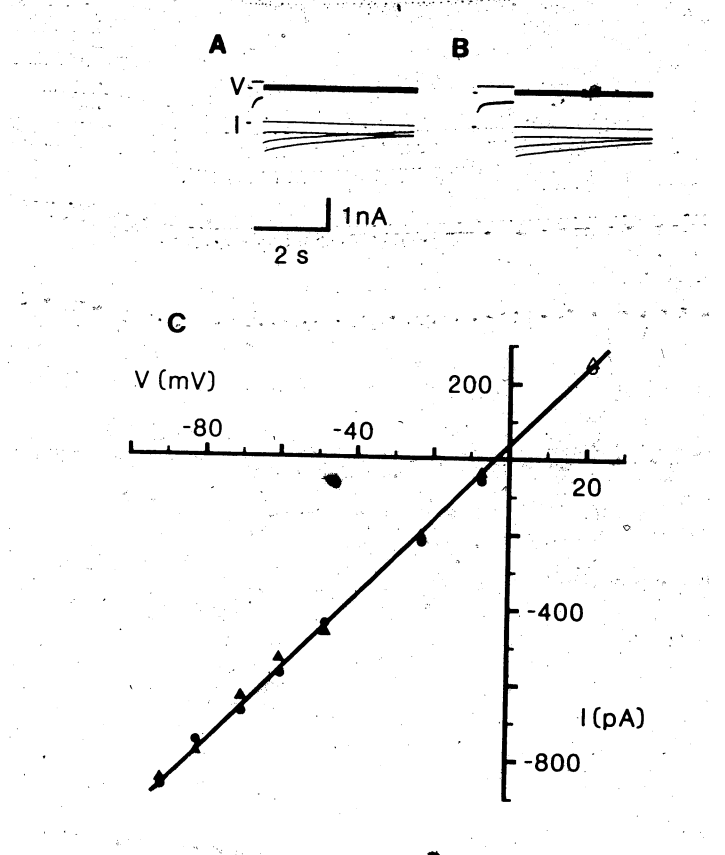




The reversal potential for tail currents was determined following depolarizing voltage steps of varying durations. The expectation was that if ion concentration gradients were changing significantly during current flow, then a shift in the reversal potential would be observed when short and long duration pulses were compared (Pigure-31). For the experishown, the reversal potential did not shift when the conditioning pulse was lengthened from 300 to 1000 ms. Similar experiments, with depolarizing conditioning potentials varying from 200 ms to 3 s, also did not show a shift in the reversal potential (n = 7). Therefore, it did not appear that the effects of changes in ion concentration gradients were causing the slow tail currents. It should be pointed out that tail currents had slow time courses even for depolarizing pulses to the reversal potential (see step to -1 mV in Figure 15). At the reversal potential, should be no net ion movement and consequently, no accumulation or depletion. In conclusion, the slow tail currents did not appear to be artifacts and must have reflected the closing behavior of the Cl channels.

Tail currents were fitted with the sum of two exponentials: a major component with amplitude A_2 (pA) and time constant τ_2 (ms) and a minor component with amplitude A, and time constant τ_1 (ms). At all potentials, the time constant was first determined for the slow, major component by fitting an exponential to the later data points and then peeling this fit away from the total current to reveal the

Figure 31. Reversal potential of tail currents for different duration test pulses. Instantaneous current-voltage plots were constructed according to the protocol in Figure 22. Depolarizing voltage step was to 21 mV from a holding potential of -70 mV. Duration of depolarizing voltage step was 300 ms in Part A and 1 s in Part B. Current-voltage plot of the data showed that the points fell along the same line with a reversal potential of -4 mV (C). Tail current magnitudes for 300 ms test pulse (filled triangles) and for 1 s pulse (filled circles). Steady-state current magnitude for the 300 ms pulse (triangle) and for the 1 s (circle). Slope of the line gave a conductance of 9.3 nS. Diameter of myoball was 30 μm which gave a unit conductance of 3.3 mS/cm². External solution was normal containing 10⁻⁷ M TTX and 1.0 mM Cd²⁺. Pipette solution was K'-free. Seal resistance was 10 G Ω . R. was 6 M Ω . R_{in} of pipette was 1 M Ω .

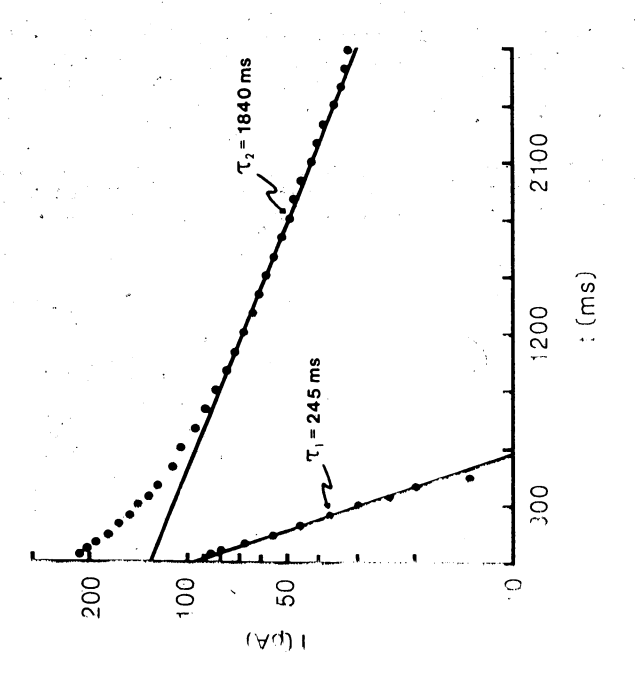


faster exponential (Figure 32). τ_2 was 8-15 times slower than τ_1 and both time constants were slightly longer at more positive potentials. For example, in one myoball, τ_1 increased from 200 ms at -90 mV to 260 ms at -60 mV. τ_2 increased from 1.74 s at -90 mV to 2.64 s at -60 mV. More time constant values are given in a later figure.

The presence of the extremely slow tail current compoment suggested that the channels entered a long-lived open state from which they returned very slowly to the closed states. The time constant for this transition at a more positive membrane potential was obtained from the following experiment. It was noted that the relative amplitude of the fast component of the tail currents depended on the duration of the conditioning pulse (Figure 33). As the depolarizing voltage step was lengthened, the amplitude of the fast tail component became smaller while the amplitude of the slow tail component became larger (Table 5). With short duration pulses (80 ms), the fast component amplitude constituted 35% of the total current while for long duration pulses (1 s), its contribution fell to about 15%. These values were plotted as a function of depolarizing test pulse duration on semilogrithmic paper (Figure 34). Data points were well fit with a single exponential with a time constant of 700 ms. This then, would be the time constant (τ_2) for the transi tion from the short to the long-lived open state.

Figure 35 shows the two times constants (τ_1 and τ_2) determined from activation and deactivation of the currents.

Figure 32. Semilogrithmic plot of tail current. Tail current following a depolarization to 38 mV was plotted as a function of time on semilog paper. Repolarization potential was -70 mV (same trace as in Figure 21). The major exponential had a time constant (τ_2) of 1840 ms and an amplitude (A_2) of 131 pA. The faster exponential is the difference between the total current and the straight line fit to the late portion of the current. The minor exponential had a time constant (τ_1) of 245 ms and an amplitude (A_1) of 100 pA).



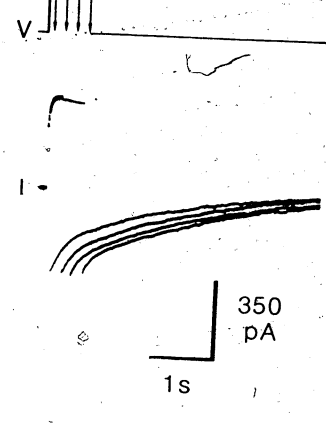


Figure 33. Effect of test pulse duration on tail currents. Depolarizing test pulse duration was varied as indicated (V). Test pulse was to 22 mV from a holding potential of 70 mV. Four current responses are superimposed (80, 240, 400, 600 ms). External solution was normal containing 10 M TTX and 1.0 mM (A). Pipette solution was 60 Cl. Stall resistance van 1 CO p. war p MO input resistance of the

Table 5. Effect of depolarizing test pulse duration on tail currents.

Test pulse was to 22 mV from a holding potential of -70 mV. Duration was increased from 80 to 1,000 msec. Tail currents were plotted on semilog paper and the two exponentials separated (see Figure 32). Data from same myoball as in Figure 33.

Duration (msec)	A ₁ (pA)	Λ ₂ (pA)	(A ₁ + A ₂) (_F Λ)	$A_1/(A_1 + A_2)$
80	141	2.72	413	0.35
120	139	295	434	0.32
160	129	317	1446	0.29
200	121	311	432	0.28
240	111	300	411	0.27
400	88	319	407	0 22
600	66	314	380	0 17
1 000	eς	5 1 1	16.1	1

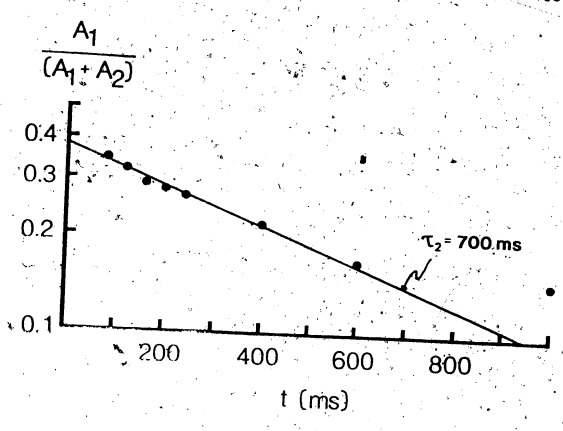
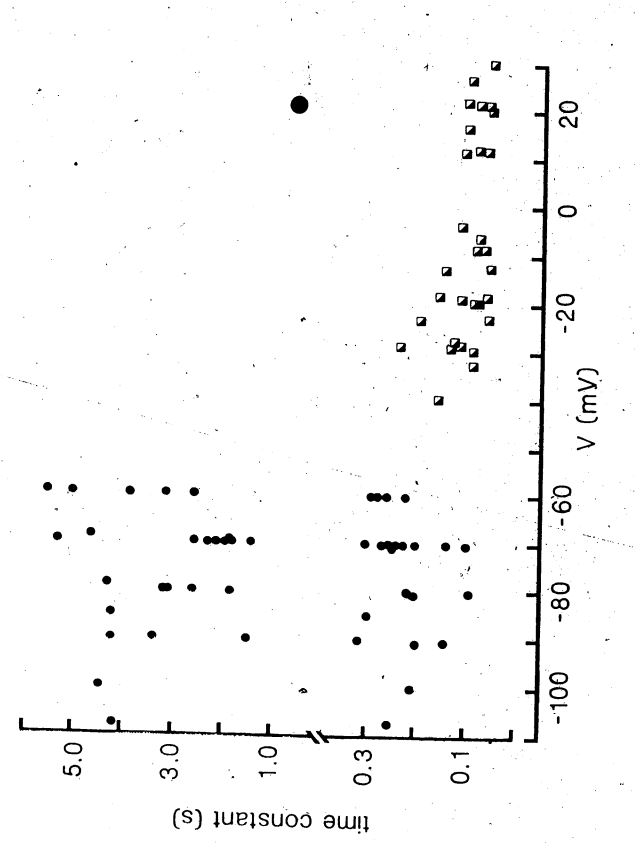


Figure 34. Semilogrithmic plot of tail current amplitude as a function of test pulse duration. $A_1/(A_1+A_2)$ was plotted semilogrithmically versus duration of the depolarizing test

Figure 35. Time constants as a function of voltage. Time constants obtained from activation and deactivation of Cl-currents are plotted versus membrane potential. Tail currents were fit with the sum of two exponentials as shown in Figure 32. Time constants from tail currents are represented by small circles. Time constant represented by the large circle is from Figure 34. Time constants obtained from activation of currents are represented by half-filled squares (data same as Figure 31). Note that the ordinate is broken.



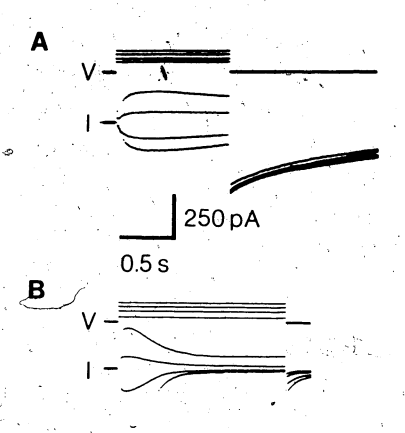


The other time constant (τ_0) obtained from activation is not shown (see Figure 30). In all, there were three measurable time constants.

Membrane currents in isolated myoballs

Chloride currents: Cl currents were recorded from myoballs that had formed spontaneously from enzymatically dissociated muscle fibers. Cl currents were found in myoballs obtained from 15 to 21 day embryos. The currents reversed at the potential predicted for Cl and showed the same kinetic properties as the currents recorded from myoballs grown in tissue culture (Figure 36A). In some myoballs, the currents declined under maintained depolarization (Figure 36B). This decline in current was also observed by Fukuda et al (1976b). The declining phase of the current could be fitted with a single exponential. Time constants for the decline were longer at positive than at negative membrane potentials. For example, in one myoball, current declined with a time constant of 410 ms for a step to -45 mV. For steps to more positive potentials, the time constants became longer and for a step to 52 mV, the time constant was 1400 ms. This behavior is opposite to that exhibited by currents which have voltage-dependent vation, i.e. the rate of decline is faster at more depolarized potentials (Hodgkin & Huxley, 1952b). Also, a plot of current magnitude versus rate of decline was nearly linear. i.e. the larger the current, the faster it declined. These

Figure 36. Chloride currents from isolated myoballs. Cl-currents were recorded under voltage clamp from a myoball obtained from 19 day embryonic intercostal muscles. Part A shows the currents recorded during step depolarizations from a holding potential of -60 mV. Four traces are superimposed. Diameter of myoball was 32 μ m. Peak current density was 7 μ A/cm². Records in B were obtained from a myoball isolated from 20 day intercostal muscles. Diameter was 27 μ m. Peak inward current density was 31 μ A/cm². External solution was normal containing 10-7 M TTX and 1.0 mM Cd². Pipette solution was K*-free.



____350 pA 2 s observations suggested that the decline of the currents resulted from ion accumulation and/or depletion. Perhaps some of the Cl channels may be located in a place where they see a restricted space, e.g. the transverse tubular system.

Other currents: Myoballs obtained from muscles of a particular age showed great variability with regard to the type and density of membrane currents. Hence, only very general conclusions could be made about the development of excitability. At the earliest ages studied, 13-14 days, most of the myoballs had no voltage- and time-dependent currents. Outward currents were found in a small percentage of myoballs. Figure 37 shows outward currents recorded from a myoball obtained from a 16 day muscle. These types of currents were absent when the pipette solution contained TEA and were presumably K currents.

Small, TTX-sensitive currents were found in myoballs obtained from 17 to 21 day muscles and were probably carried interesting observation was made for this by Na . Αn Ιf the myoballs were kept overnight in tissue culture, the current density increased about 25 fold. For example, for fresh 17 day myoballs the inward current density averaged 13 μ A/cm². After overnight incubation, current density had increased to around 300 $\mu A/cm^2$. This value was much greater than for freshly isolated 18 day myoballs (Figure 38 & 39). It was as if the expression of Na $^{+}$ channels was being suppressed by something in the embryo.

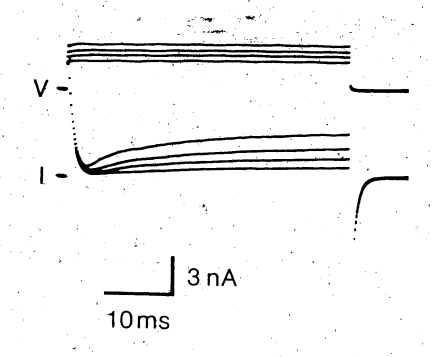


Figure 37. Outward currents from isolated myoball. Currents were recorded under voltage clamp. Depolarizing voltage steps were applied from a holding potential of -70 mV. Four traces are superimposed. Myoball was obtained from 16 day embryonic intercostal muscles. External and pipette solutions were normal.

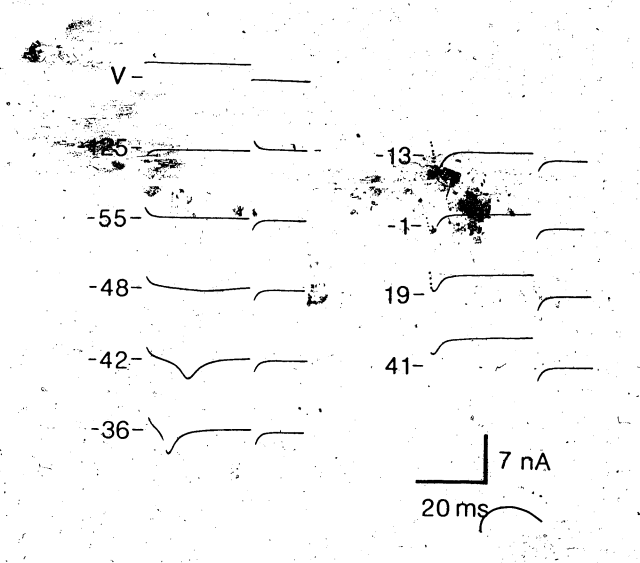
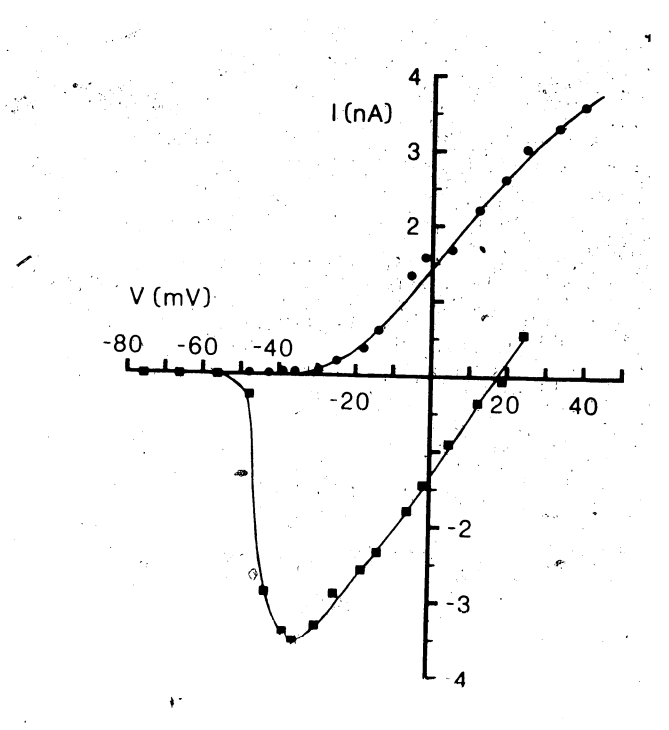


Figure 38. Inward and outward currents from isolated myoballs. Currents were recorded under voltage clamp from a myoball that had been obtained from 18 day intercostal muscles and kept overnight in tissue culture. Depolarizing test potential is noted beside each trace. Holding potential was -90 mV. External and pipette solutions were normal. Seal resistance was 5 G Ω . R. was 17 M Ω . Input resistance of pipette was 5 M Ω . Traces 7, 8 and 9 were retouched.

Figure 39. Current-voltage plot of inward and outward currents. Plot of peak inward currents (squares) and steady-state current (circles) as a function of depolarizing test potential. Diameter of myoball was 27.5 μ m. Peak inward current density was 14.9 mA/cm² and outward current density was 15.4 mA/cm² (measured at 40 mV). Data from currents shown in Figure 38 and from additional steps.



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Sherman et al. (1982) have also observed an increase, although not as large, in Nat channel density following denervation of developing rat muscle. In addition, it has been reported that slow-tonic muscle of the frog, which normally does not have Nat channels, expresses them following denervation (Cull-Candy, Miledi & Uchitel, 1980).

In summary, it appeared that the outward currents were the first to appear during development. The invaded currents appeared next and then the Mai currents.

D. DISCUSSION

Chloride currents in embryonic chick muscle

The whole cell patch clamp technique (Hamil' et al., 1981) was found to provide adequate voltage control of the surface membranes of myoballs obtained from chick skeletal muscles. Slow membrane currents were recorded from myoballs grown in tissue culture (Figure 15 and from myoballs obtained from muscle that had developed in the (Figure 36). The currents were shown to be accorded by the reversal potential for the arrents varied with (1) concentrations according to the values predicted by the Dernot equation (Figure 18). Also, the currents were blocked by the stilbene derivatives. SITS and DIDS (Figure 19) which have been shown to block (1) cormeability in cariety of cell types (Enauf the Doths'sia, 1971) Usugha & Fong (2) figure 19) the

openings of 'Cl channels', as yet hypothetical macro-molecules embedded in the membrane which permit transmembrane Cl movements.

The Cl currents were shown to produce a long duration action potential (Figure 14). Depolarization of the membrane potential was brought about by inward Cl currents, i.e. Clions were moving out of the myoballs. In order for a Claction potential to be generated, the equilibrium potential for Cl must be more positive than the resting membrane potential. Therefore, the intracellular Cl concentration must be higher than that which would occur as a result passive equilibration. This suggests that the embryonic chick fibers have an inward Cl pump Inward Cl pumps have shown to exist in squid axon (Keynes, 1963; Russell, 1979) and suggested to exist in some mammalian muscles LDulhunty, 1978). Outward Cl rumps have been demonstrated in Aplysia neurones (Russell & Prown, 1972: Russell, 1978), in Farnacle skeletal mustle (Possell & Brodwick, 1979 and in cat moto eurones (Tux, 1071; Idinas, Poker & 1774) The freernal rotential of the Cl action potentials was around 20 to 25 mm in chick muscle fibers (Figure 2) (Kano, 1975). From the Nernst equition, this gives an intracelly far Cl. ,concentration of about 60 mM. The concentration was probably not a high in cultured chick muscle as the riversal notential of the action potential was more requires (50 mm) . This may be the rest that the mem-. . 1

injected intracellularly in order to elicit Cl action potentials in cultured muscle (Fukuda, 1974; Spector & Prives, 1977).

In cultured chick myotubes, it has been observed that the depolarization phase of the Cl action potential coincided with the development of contractures, while the repolarization phase coincided with relaxation (Spector & Prives, 1977). In developing p.l.d. muscles of the chick, extremely long duration contractions have been recorded in response to 1 ms extracellular stimulation (Peiser & Stokes, 1982). The time ourse of these unusual contractions and the ages at which they occurred strongly suggest that the Cl action potential was involved. Thus, the Cl action potential appears to be involved in coupling in the emission (bees.

Commercian with other chloride channels

studied in detail Of the other Of channels have centraled in detail Of the other Of channels, only the channel from Identifications. Identification of channels were incomporated into plans; the channels to disperse and todical under voltage of all channels of dust need to be about 15 per (while a Mill of 1972; from an the famels channel tabories a systemel; complime as the famels displayed multification of the complime as the famels displayed multification of the complimentations.

channels showed sigmoidal dependence of conductance on voltage (Figure 24). Conductance was low (channels closed) at negative potentials and high (channels open) at positive potentials (White & Miller, 1979). An exact correlation of conductance with absolute membrane potential is not possible since the position of the conductance curve along the voltage axis was affected by the phospholipid composition of the bilayers. The voltage sensitivity of chick and Torpedo channels was very similar. Conductance changed e-fold for an 11 mV depolarization as compared to an 8 mV change determined for chick Cl channels (Figure 26). Both conductances were blocked by SITS and DIDS (Figure 19) (White & Miller, 1979) and by SCN (Figure 19) (White & Miller, 1979). The kinetics of the macroscopic currents were not studied in detail for the Torpedo Cl channels. However, the currents did not show any inactivation which was the case for the Cl currents in this study (Figure 15, 20). The turn on and the turn off of the macroscopic currents followed single exponential time courses. The rise times appeared to be much longer, on the order of tens of seconds, those seen for chick Cl currents (Figure 15). Also, chick Cl currents clearly showed two exponentials during Promission (Figure 32) while Torpedo Cl currents did not.

Interestingly, the kinetics of the channel-forming profess, voltage dependent anion conductance (VDAC), from outer mitochandrial membranes (Colombini, 1979) are reministrated of the inetic of the currents studied here. The

kinetics of the macroscopic VDAC currents have not been studied in detail, but the following similarities were apparent (Colombini, 1979). Turn-on of the macroscopic current appeared to follow a single exponential time course with the rise time being very close to the 50-100 ms range found for chick Cl currents. Closing of the VDAC channels was very slow, on the order of seconds. There also appeared to be two exponential phases. This is very similar to the properties found for the chick Cl currents (Figure 32). However, the similarity between the two channels stops there. Single channel conductance for VDAC channels has been reported to be about 500 pS (Colombini, 1979). It unlikely that the conductance of chick Cl. channels is that high as the macroscopic currents would have been very noisy. Also, the dependence of conductance on voltage for the two channels is radically different. VDAC channels have been shown to open only over a very narrow range of potentials around 0 mV (Doring & Colombini, 1984)

Chloride current kinetics

The kinetics of the Cl currents were more complex than could be described by a simple Hodgkin Huxley kinetic model (1952b). According to their model, tail currents measured at very negative potentials should display a single exponential component. For chick Cl currents, however, there were clearly two exponential components even at very negative potential (Figure 32). An alternative to the U dakin Huxley

model is to suppose that the channels must pass through a sequence of kinetically distinct states. Since three distinct time constants were measured (Figure 30, 35), there must be at least four kinetic states (Colquhoun & Hawkes, 1983). The following kinetic model was developed:

$$C_2 = C_1 = O_1 = O_2$$

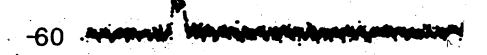
In this scheme, C_2 and C_1 represent closed states of the channel and O_1 and O_2 represent open states. The presence of the second closed state (C_2) was needed to account for the delay in activation (Figure 28). Although two closed states are indicated, the exact number may be greater than 2. The second open state (O_2) was needed to account for the slow exponential tail current component (Figure 21). Thus, during a depolarizing pulse, channels would enter the open states sequentially. When the membrane potential is repolarized, C1 current would decline rapidly as channels proceeded from O_1 to C_1 . But a slowly declining phase of current would also be present as channels in O_2 returned slowly to the closed states by way of O_1 .

Evidence to support the hypothesis that the channels have two kinetically distinct open states can be obtained by observing the kinetics of single channel currents. The amount of time spent in a given kinetic state (dwell time) has been shown to be exponentially distributed (cf Horn & Lange, 1983). Thus, it is predicted that the open time histograms of the Cl channels could be fitted with the sum of two exponentials. Attempts were made to record single

channel C1 currents in outside-out membrane patches. Single channel events could be recorded (Figure 40); however, no C1 channels with the appropriate voltage-dependent behavior were consistently observed. It was quite possible that the conductance was too small to be detected with the noise level of my set-up (any channel under 10 pS would have been difficult to detect). Or, perhaps the channel did not survive in outside-out membrane patches. Some channels, for example Ca² channels (Hagiwara & Byerly, 1981), require intracellular components for activity and do not survive long in excised membrane patches.

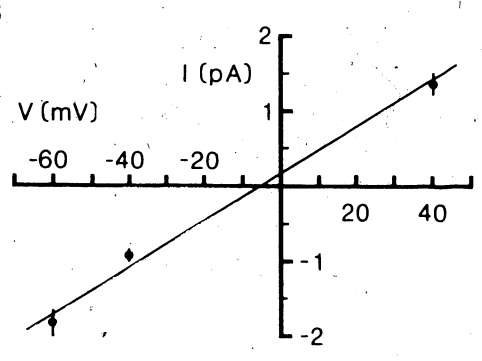
Figure 40. Single channel events. Single channel currents were recorded in outside-out membrane patches (see Figure 5) from cultured chick myoballs. Channel openings are displayed upwards (A). Holding potentials are indicated beside each current trace. External solution was normal containing 10⁻⁷ M TTX and 1.0 mM Cd²⁺. Pipette solution was K⁺-free. Current-voltage plot gave a single channel conductance of 31 pS (B). Measured reversal potential was -5 mV. Values are means ± S.D. for 15 channel openings.

A



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B



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